

Constraints on emissions of carbon monoxide, methane, and a suite of hydrocarbons in the Colorado Front Range using observations of 14CO2

B. W. LaFranchi, G. Petron, J. B. Miller, S. J. Lehman, A. E. Andrews, E. J. Dlugokencky, B. R. Miller, S. A. Montzka, B. Hall, W. Neff, C. Sweeney, J. Turnbull, D. E. Wolfe, P. P. Tans, K. R. Gurney, T. P. Guilderson

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1 Constraints on emissions of carbon monoxide, methane, and a suite of

- 2 hydrocarbons in the Colorado Front Range using observations of ¹⁴CO₂
- 3 B.W. LaFranchi*¹, G. Pétron^{2,3}, J.B. Miller^{2,3}, S.J. Lehman⁴, A.E. Andrews², E. Dlugokencky², B.R. Miller^{2,3},
- 4 S.A. Montzka², B. Hall^{2,3}, W. Neff^{3,5}, C. Sweeney², J.C. Turnbull⁶, D.E. Wolfe⁵, P.P. Tans², K.R. Gurney⁷, T.P.
- 5 Guilderson¹

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- Center for Accelerator Mass Spectrometry (CAMS), Lawrence Livermore National Laboratory,
 Livermore, CO 94550 USA
- 8 2) Global Monitoring Division (GMD), NOAA Earth Systems Research Laboratory, Boulder, CO 80305
 9 USA
- 3) Cooperative Institute for Research in Environmental Sciences (CIRES), University of Colorado,
 Boulder CO 80309 USA
- 12 4) Institute for Arctic and Alpine Research (INSTAAR), University of Colorado, Boulder 80305 USA
- 13 5) Physical Sciences Division, NOAA Earth Systems Research Laboratory, Boulder, CO 80305 USA
- 14 6) National Isotope Centre, GNS Science, Lower Hutt 5040 New Zealand
- 15 7) School of Life Sciences, Arizona State University, Tempe, AZ 85287 USA
- *corresponding author: lafranchi2@llnl.gov, (925)423-4555

Abstract

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Atmospheric radiocarbon (14C) represents an important observational constraint on emissions of fossilfuel derived carbon into the atmosphere due to the absence of ¹⁴C in fossil fuel reservoirs. The high sensitivity and precision that accelerator mass spectrometry (AMS) affords in atmospheric ¹⁴C analysis has greatly increased the potential for using such measurements to evaluate bottom-up emissions inventories of fossil fuel CO₂ (CO₂ff), as well as those for other co-emitted species. Here we use observations of ¹⁴CO₂ and a series of hydrocarbons and combustion tracers from discrete air samples collected between June 2009 and September 2010 at the National Oceanic and Atmospheric Administration Boulder Atmospheric Observatory (BAO; Lat: 40.050°N, Lon: 105.004°W) to derive emission ratios of each species to CO₂ff. From these emission ratios, we estimate emissions of these species by using the Vulcan CO₂ff high resolution data product as a reference. The species considered in this analysis are carbon monoxide (CO), methane (CH₄), acetylene (C₂H₂), benzene (C₆H₆), and C₃-C₅ alkanes. Comparisons of top-down emissions estimates are made to existing inventories of these species for Denver and adjacent counties, as well as to previous efforts to estimate emissions from atmospheric observations over the same area. We find that CO is overestimated in the 2008 National Emissions Inventory (NEI 2008) by a factor of ~2. A close evaluation of the inventory suggests that the ratio of CO emitted per unit fuel burned from on-road gasoline vehicles is likely over-estimated by a factor of 2.5. The results also suggest that while the oil and gas sector is the largest contributor to the CH₄ signal in air arriving from the north and east, it is very likely that other sources, including agricultural sources, contribute to this signal and must be accounted for when attributing these signals to oil and gas industry activity from a top-down perspective. Our results are consistent with ~60% of the total CH₄ emissions from regions to the north and east of the BAO tower stemming from the oil and gas industry, equating to ~70 Gg yr⁻¹ or ~1.7% of total natural gas production in the region.

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1. Introduction

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Radiocarbon in CO₂ (¹⁴CO₂) is a powerful tracer that provides the least biased and most direct means to 66 67 observe fossil fuel derived CO₂ in the atmosphere (Zondervan and Meijer, 1996; Hsueh et al., 2007; Levin 68 and Karstens, 2007; Turnbull et al., 2009; Van der Laan et al., 2010). Fossil fuels are completely devoid of 69 14 C, as is the CO₂ resulting from its combustion, because the half life of 14 C is short ($^{\sim}$ 5700 years (Godwin, 1962)) in relation to the residence times of carbon in fossil reservoirs, where no additional ¹⁴C 70 71 production occurs. Since all other sources of CO2 to atmosphere stem from carbon reservoirs that are in 72 near equilibrium with the isotopic composition of the atmosphere itself, the atmosphere exhibits gradients in ¹⁴CO₂ that can be quantitatively traced to addition of CO₂ from fossil fuel combustion 73 (Turnbull et al., 2007; Graven et al., 2009; Levin et al., 2010). 74

Prior to nuclear weapons testing, which artificially increased the ¹⁴CO₂ content of the atmosphere, the rise in atmospheric CO₂ resulting from fossil fuel combustion could be observed on global scales as a decrease in ¹⁴CO₂, widely known as the Seuss effect (Suess, 1955). While ¹⁴CO₂ is produced naturally in the upper atmosphere from cosmogenic radiation, the abundance of ¹⁴CO₂ in the modern atmosphere was strongly impacted by above ground nuclear testing that occurred in the middle part of the 20th century. Since the atmospheric nuclear weapons test ban was put in place, the decrease in ¹⁴CO₂, which has been observed at a number of global background monitoring sites (Levin and Kromer, 2004; Turnbull et al., 2007; Currie et al., 2011; Graven et al., 2012a, b), has been influenced primarily by the equilibration of atmospheric ¹⁴CO₂ with the oceanic and terrestrial carbon reservoirs. In recent years, however, the atmospheric decline has been dominated increasingly by isotopic dilution due to the Seuss effect, as fossil fuel combustion increases and as the atmosphere, ocean, and terrestrial carbon reservoirs approach equilibrium with the "bomb spike". On regional scales, locally emitted CO₂ from fossil fuel combustion can be detected as a depletion of ¹⁴C:¹²C relative to background air. These observed gradients result from what we define as "recently added" fossil-fuel CO₂ (CO₂ff).

Observations of ¹⁴CO₂ are of great interest, not only for the evaluation of fossil CO₂ emissions inventories, but also as a means to better understand emissions of a range of trace gases associated with the combustion of fossil fuels (Turnbull et al., 2011; Miller et al., 2012). Bottom-up inventories of these trace gases carry significant uncertainties because of the difficulty in quantifying the relationship between the mass of fuel consumed and the mass of trace gas emitted. Emissions of by-products, including species such as carbon monoxide (CO), methane (CH₄), acetylene (C_2H_2), and benzene (C_6H_6) depend on a number of variables including fuel type, combustion temperature, the extent of tail-pipe or flue-stack "scrubbing", and oxidant-to-fuel ratio. For example, it has long been known from observations that the National Emissions Inventory (NEI) appears to over-estimate observed anthropogenic emissions of CO in the United States (Parrish, 2006; Hudman et al., 2008; Miller et al., 2008; Miller et al., 2012), though the reason remains unclear. Further, there are a number of industrial activities that lead to noncombustion emissions of gases impacting air quality and climate from leaks in transmission lines, venting of storage tanks, and other processes, in which case, quantifying emissions based on readily available fuel use statistics is difficult or not possible. In contrast, the amount of CO₂ emitted per unit of fuel combusted can be derived stoichiometrically with relatively high accuracy. Accordingly, the bottom-up inventory of fossil fuel derived CO₂ in the United States (e.g. EIA, 2012) and in most developed countries

is thought to be relatively reliable. Estimates of annual fossil CO_2 emissions for the U.S. are thought to be reliable to ~10% (1 σ) (Gurney et al., 2011), though uncertainties are larger at smaller spatial and temporal scales.

To improve emissions estimates for these other combustion and industrial tracers, particularly those species that can affect air quality, human health, and climate, observations in the atmosphere are necessary for critically evaluating the existing bottom-up inventories. One relatively simple strategy for deriving emissions based on atmospheric observations is the use of tracer/tracer enhancement ratios in which emission ratios of two species are inferred from the ratio of the observed mole fraction enhancements (with respect to background observations) of one species to the other. For gases with lifetimes comparable to the transit times between emission and measurement, a simple photochemical age model can be used to extrapolate back from the time of the observation to derive the ratio at the time of emission (Lee et al., 2006; Warneke et al., 2007). Then, if emissions of one of the tracers are relatively well defined for the geographic area that the observations are sensitive to, emissions of the other tracer can be calculated from the inferred emission ratio. Uncertainties for this method are minimized when both tracers have long atmospheric lifetimes and slow atmospheric production rates on the time scales relevant to the source-receptor distances. A major advantage of this approach comes from its computational simplicity. Additionally, since all tracers are expected to be mixed and transported in the same way if their sources are co-located, this approach reduces the sensitivity of the analysis on uncertainties in transport and boundary layer calculations.

The very long atmospheric chemical lifetime of CO₂ would be ideal for use in these tracer/tracer approaches, but large uncertainties in its biogenic sources and sinks can complicate its use in inferring emissions of other fossil fuel combustion tracers (e.g. Miller et al., 2012). Thus, to take advantage of the photochemical stability of CO₂ and the availability of relatively accurate fossil fuel CO₂ emissions inventories measurements of ¹⁴CO₂ must be used to isolate the fossil fuel contribution to the observed CO₂. Here we describe observations of ¹⁴CO₂ and other trace gases made between June 2009 and September 2010 at the Boulder Atmospheric Observatory (BAO), a 300 m tall tower located 35 km north of Denver, CO (Lat 40.05N, Lon 105.01W). BAO is one of 9 towers in the NOAA Earth System Research Laboratory Global Monitoring Division (NOAA-GMD, hereafter) tall tower network (Andrews et al., submitted). It is one of 7 towers in the network that is monitoring CO₂ and CO continuously and collecting air samples daily for multiple species analysis and one of 6 that also measures ¹⁴CO₂. The observations presented here represent the first report of ¹⁴CO₂ observations from this network. In this study, we use ¹⁴CO₂ to derive CO₂ff mole fractions for estimating emission ratios and emissions of a number of important trace gases being transported to the site from the Denver metro region and from the extensive oil and gas drilling operations and cattle feedlots in Weld County, to the northeast.

This study builds on a previous effort to characterize emissions of volatile organic compounds (VOCs) and CH₄ from oil and gas production and drilling operations in Weld County using both bottom-up and top-down approaches for 2008 (Pétron et al., 2012). We will refer to this prior study as the Colorado Front Range Pilot Study (CFRPS, hereafter), in which the authors made use of observations at BAO in combination with those from a mobile platform to determine emission signatures of individual methane sources, including oil and gas wells, natural gas processing plants, condensate storage tanks, landfills,

cattle feed-lots, and waste water treatment plants. A crucial finding of the CFRPS is that trace gas concentrations measured at BAO are influenced most substantially by two different source regions: oil and gas fields in Weld County (from a region known as the Denver Julesberg Basin, or DJB) and urbantype emissions from the Denver metro region. They found that air arriving at BAO from the northeast exhibits strong enhancements in alkanes, including methane, resembling enhancements (based on tracer/tracer ratios) similar to those sampled on the mobile platform within the DJB. These results suggested that oil and gas operations are the dominant emitters of alkanes and methane in the region and that the observations at BAO provide reliable constraints on emissions from these activities.

The CFRPS estimated emissions from the DJB using a tracer/tracer approach with constraints set by bottom-up statistical data on methane and propane emissions from a subset of condensate tanks in the region and a survey of raw natural gas composition. The confidence in these estimates is ultimately limited by uncertainties in the average composition of gas released from a representative sub-set of condensate tanks and oil and gas wells exhibiting a wide range of compositions. In a separate evaluation of the data analyzed in the CFRPS, Levi (2012) (denoted as L12, hereafter) using a slightly different approach making use of n-butane data in addition to methane and propane, estimated CH₄ emissions that are substantially lower than the CFRPS estimates and are more in line with the bottom-up estimates. Here, we add to the characterization of trace gases emissions in the DJB, as well as in the Denver metro region, by using radiocarbon-derived CO₂ff observations as an improved constraint that is independent of the assumptions made in these two prior studies. The primary advantage of the approach taken here is that by using the Vulcan fossil fuel CO₂ data product (Gurney et al., 2009), which is reliable nationwide to within 20% at the county level on annual time-scales, our confidence in the reference emissions estimate is greatly improved. Further, as we will show, CO₂ff exhibits strong correlations with a variety of trace gases in the region, both from combustion and non-combustion sources, thus allowing for the evaluation of emissions from a range of different source-types.

2. Methods

2.1. Site Description

The BAO tower is located 25 km east-northeast of Boulder and 35 km north of Denver (40.05° N, 105.01° W). The base of the tower is at 1584 m above sea level (a.s.l.). As shown in Figure 1, BAO is located at the southwestern edge of the DJB where a very large and dense network of oil and gas wells exists. Since late 2007 NOAA-GMD has been collecting discrete air samples approximately daily from 300 m. The air is collected in glass flasks and analyzed at NOAA-GMD for a suite of ~50 trace gases and then circulated to the University of Colorado's Institute of Arctic and Alpine Research (INSTAAR) for stable isotope measurements in CO_2 and CH_4 and preparation for $^{14}CO_2$ measurement. The Center for Accelerator Mass Spectrometry (CAMS) at Lawrence Livermore National Laboratory (LLNL), which performed the $^{14}CO_2$ measurements reported here, has participated in the NOAA-GMD $^{14}CO_2$ discrete air sample measurement program since 2009. This study focuses on data collected between June 2009 and September 2010, over which time 145 samples were analyzed for $^{14}CO_2$. More information on this site and the entire tall tower network can be found at: http://www.esrl.noaa.gov/gmd/ccgg/towers/.

- 184 Standard meteorological measurements are also made continuously at several levels on the tower by
- the NOAA ESRL Physical Sciences Division, including wind speed and direction, relative humidity, and
- temperature. We categorize each observation in our analysis according to wind direction in order to
- 187 facilitate a discussion of two distinct emission source regions: the oil and gas industrial region to the
- north and east and the Denver metro region to the south. To do this, we define three wind sectors,
- consistent with those defined in the CFRPS: N/E (345° to 120°), S (120° to 240°), and W (240° to 345°).
- 190 These wind sectors are illustrated in Fig. 1. Wind direction for each sample is determined using the
- mean wind direction over the 30 minutes immediately prior to sampling. Samples with mean wind
- speeds lower than 2.5 m/s over this time period are removed from any sector-specific analysis in this
- 193 study.
- 194 To define isotopic and mole fractions of trace gases in background air, measurements from two
- additional sites were used. For ¹⁴CO₂, CO₂, CO, and CH₄, we used weekly measurements from Niwot
- Ridge, CO (sitecode NWR, 40.05 °N, 105.63 °W, 3526 m a.s.l.), a site in the alpine tundra with strong
- 197 westerly winds that only rarely required filtering of samples influenced by pollution from the Denver
- metro area (Turnbull et al., 2007). For other gases, including acetylene, benzene, and the C₃-C₅ alkanes
- we used weekly to fortnightly samples collected in the free troposphere from flights at a nearby location
- 200 (3000 to 4000 m a.s.l. above Briggsdale Colorado; sitecode CAR, 40.37 °N, 104.30 °W, ground elevation
- 201 ~1700 m a.s.l.).

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2.2. Flask Sampling

- 203 Discrete whole air samples are collected daily from the BAO tall tower (at the 300 m level) using
- 204 Programmable Flask Packages (PFPs) connected to a Programmable Compressor Package (PCP) capable
- of delivering 15 standard L min⁻¹ (Andrews et al., submitted). Each PFP contains 12 cylindrical
- borosilicate glass flasks (0.7 L each). On each end of the flasks are automated glass-piston stopcocks,
- sealed with Teflon O-rings. Prior to deployment, each flask in the PFP unit is flushed with clean dry air
- and then pressurized to ~140 kPa with synthetic air containing 330 ppm CO₂ (Sweeney et al., in prep).
- Automated sampling consists of the following steps: (1) a manifold flush, (2) a flask flush, and (3)
- 210 pressurization of the flask to ~270 kPa. The entire process takes about 2 minutes. Sampled air at BAO
- 211 first passes through a drying stage (dewpoint temperature at ambient pressure of ~5 °C) prior to
- collection. Sampling is done at midday (1930 UTC) in most cases; all samples used in this analysis were
- collected within about 30 minutes of 1930 UTC. Two flasks are filled within 5 minutes of each other (~4
- standard liters) for analysis of the standard suite of trace gases (described below), with enough gas
- remaining for analysis of ¹⁴CO₂, which typically requires 0.4 to 0.5 mg C for high precision analysis.

2.3. Flask Analysis

- Each flask pair is analyzed at NOAA–GMD for CO₂, CO, CH₄, SF₆, H₂, N₂O, and a suite of halocarbons and
- 218 hydrocarbons. Stable isotopes of CO_2 ($\delta^{13}C$ and $\delta^{18}O$) are analyzed at the INSTAAR Stable Isotope
- Laboratory (Vaughn et al., 2004). In this study, we use measurements of CO₂, CO, CH₄, acetylene (C₂H₂),
- benzene (C_6H_6), propane (C_3H_8), n-butane ($n-C_4H_{10}$), n-pentane ($n-C_5H_{12}$), and i-pentane ($i-C_5H_{12}$). We also
- use δ^{13} C in CO₂ in the calculation of Δ^{14} C, according to methods described by Stuiver and Polach (1977).

- 222 Dry air mole fractions of CO₂, CH₄, and CO were measured on one of two nearly-identical custom
- automated analytical systems. These systems consist of custom-made gas inlet systems, gas-specific
- analyzers, and system-control software. During this project, each system used a different technique to
- measure CO. One used a Reduction Gas Analyzer, where CO is separated from air by gas
- 226 chromatography, then passed through a heated bed of HgO producing Hg before it is detected by
- resonance absorption (Novelli et al., 1998). The second is Vacuum UV Resonance Fluorescence (VURF),
- 228 where CO is detected by fluorescence at ~150 nm. Both techniques are calibrated against the same
- 229 standard scale, and uncertainties (68% confidence interval) are ~1 ppb for the VURF and ~2 ppb for the
- 230 RGA. Long-term comparison of the two systems show the RGA and VURF agree within ~ 1 ppb. CH₄ was
- 231 measured by gas chromatography (GC) with flame ionization detection with an uncertainty of ~1.4 ppb
- 232 (Dlugokencky et al., 1994). A non-dispersive infrared analyzer is used for CO₂ with an uncertainty
- 233 <0.1ppm (Conway et al., 1994).
- The hydrocarbons (C_2H_2 , benzene, and C_3 - C_5 alkanes) are measured using a GC/mass spectrometric
- 235 technique, with cryogenic pre-concentration (Montzka et al., 1993). Relative measurement uncertainties
- for the hydrocarbons considered in this study vary depending on mole fraction, with lower relative
- 237 uncertainties estimated for samples with lower mole fractions (down to 10 ppt). For mole fractions
- between 10 ppt and 1 ppb, uncertainties (1 σ) are 5% for i-C₅H₁₂, n-C₅H₁₂, and C₆H₆, and 15% for C₃H₈,
- C_2H_2 and $n-C_4H_{10}$. At higher atmospheric mole fractions the uncertainties are larger: by a factor of two
- 240 for mole fractions between 1 ppb and 20 ppb, and a factor of three for mole fractions between 20 ppb
- and 50 ppb. Only C₃H₈ was observed at mole fractions greater than 50 ppb during the study period;
- these 2 samples were removed from this analysis. The average relative uncertainties for individual
- samples are 5-8% for $i-C_5H_{12}$, $n-C_5H_{12}$, and C_6H_6 and 15-30% for C_3H_8 , C_2H_2 and $n-C_4H_{10}$ during the study
- period. Measurement repeatability (1σ) is generally <2% for compounds present at mole fractions > 10
- ppt. For C₂H₂ and C₃H₈, the most volatile of these compounds, repeatability was somewhat poorer
- 246 during these flask analyses (approx. -25% and +12%).
- 247 All measurements are reported as dry air mole fractions relative to internally consistent standard scales
- 248 maintained at NOAA-GMD. We use the following abbreviations for measured dry air mole fractions:
- ppm = μ mol (trace gas) mol (dry air)⁻¹, ppb = nmol mol⁻¹, and ppt = pmol mol⁻¹. Additional details on
- 250 these methods are described at http://www.esrl.noaa.gov/gmd/ccgg/aircraft/analysis.html.

2.4. Radiocarbon Analysis

- A subset (typically 1 out of every 2 pairs) of the flask pairs are hand selected for analysis of ¹⁴CO₂. The
- selection is based on an analysis of continuous CO and CO₂ observations with the intent of selecting a
- 254 combination of both locally-impacted and background samples. High precision measurements of ¹⁴CO₂
- 255 were made by extracting CO₂ from the whole air samples using cryogenic separation, reducing the
- extracted CO₂ to graphite for atom counting via accelerator mass spectrometry (AMS). Extractions of
- authentic samples, measurement controls, and process blanks were performed at the University of
- 258 Colorado INSTAAR Laboratory for AMS Radiocarbon Preparation and Research (NSRL) using an
- automated extraction system (Turnbull et al., 2010). Graphitization and AMS analysis was done at LLNL-
- 260 CAMS. A description of the high precision methods for analysis of atmospheric samples at CAMS is given

by Graven et al. (2007). The measurements are expressed as age-corrected $\Delta^{14}CO_2$ in units of per mil (‰), calculated from the $^{14}C/^{13}C$ ratio, measured relative to NBS Oxalic Acid I (OX1) and reported relative to the absolute radiocarbon standard (1890 wood), as detailed in Stuiver and Polach (1977).

Uncertainty in these observations is assigned as the standard deviation (1σ) of a series of repeat measurements on extraction aliquots of whole air stored in high pressure cylinders. Air from two surveillance cylinders having different but near-ambient 14 C activities, identified as NWT3 and NWT4, were extracted, graphitized, and analyzed concurrent with the BAO samples across 7 different measurement "wheels" or batches. Multiple samples of NBS Oxalic Acid II (OX2, a commonly used secondary standard) were combusted, graphitized and analyzed simultaneously. Typically, in a wheel containing 25 authentic samples, 12 measurement controls and 1 process blank were analyzed. For the observations described in this study, the (1σ) repeatability (standard deviation) of NWT3 and NWT4 samples was $\pm 2.2\%$. In a small number of cases, the internal variability of repeat measurements of the same sample was larger than the repeatability of the pool of NWT samples. The larger of the two is assigned as the uncertainty for a given Δ^{14} CO2 measurement.

2.5. Calculation of CO₂ff

Recently added fossil fuel CO_2 (CO_2 ff) is defined as the local enhancement of CO_2 , with respect to an appropriate background reference site, due to fossil fuel emissions. CO_2 ff is estimated using a mass balance approach (Levin et al., 2003), in which the observed mole fraction of CO_2 (CO_2 obs) is partitioned into background CO_2 (CO_2 bkg), fossil CO_2 , and biogenic CO_2 (CO_2 bio) components. CO_2 bio is the net balance between respired CO_2 (CO_2 resp) and CO_2 taken up by photosynthesis (CO_2 photo). We further separate the respired fraction into autotrophic respiration (CO_2 auto) and heterotrophic respiration (CO_2 het) that originates from older soil carbon pools (which typically contain more bomb ^{14}C). Equations (1a) and (1b) detail this mass balance relationship, as formulated in Turnbull et al. (2006) and also described in Miller et al. (2012), with CO_2 resp separated into heterotrophic and autotrophic components. Similarly, an isotopic mass balance equation (Eq. (2)) can describe the contribution of these three end members to the total observed $\Delta^{14}C$.

$$CO_2obs = CO_2bkg + CO_2ff + CO_2bio$$
 (1a)

$$CO_2bio = CO_2auto + CO_2het - CO_2photo$$
 (1b)

$$\Delta_{obs}^{14}CO_{2}obs = \Delta_{bkg}^{14}CO_{2}bkg + \Delta_{ff}^{14}CO_{2}ff + \Delta_{bio}^{14}CO_{2}bio$$
 (2)

Since Δ^{14} C values are all normalized by their δ^{13} C values, and thus are not influenced by natural fractionation, we can assume that Δ^{14}_{photo} and Δ^{14}_{auto} are identical to Δ^{14}_{bkg} (Turnbull et al., 2006). The system of equations can then be solved for CO₂ff to give Eq. (3).

$$CO_2 ff = \left(\frac{CO_2 obs(\Delta_{obs}^{14} - \Delta_{bkg}^{14})}{(\Delta_{ff}^{14} - \Delta_{bkg}^{14})}\right) - \left(\frac{CO_2 het(\Delta_{het}^{14} - \Delta_{bkg}^{14})}{(\Delta_{ff}^{14} - \Delta_{bkg}^{14})}\right)$$
(3)

In this equation, the variables in the first term are either known (Δ^{14}_{ff} = -1000%) or can be measured. We use observations from NWR to estimate Δ^{14}_{bkg} . The background estimate is calculated by applying a

smoothing algorithm (Thoning et al., 1989) to the NWR data (a curve-fit of 3 polynomials, 4 harmonics, and added low-pass filtered residuals), after filtering out samples influenced by upslope flows carrying locally influenced air, characterized by high CO/CO_2 ratios, as in Turnbull et al. (2007). The selection of a proper background site introduces uncertainties on the order of the measurement uncertainty (~2 ‰) (Turnbull et al., 2009). The second term in Eq. 3 is a minor correction to the calculation of CO_2 ff due to heterotrophic respiration from soils, which can draw from carbon pools that are on the order of 10 years old and, thus, reflect the higher $\Delta^{14}CO_2$ in the atmosphere at the time. The magnitude of this correction can be estimated from a terrestrial ecosystem model, such as the CASA biogeochemical model (Thompson and Randerson, 1999); we follow the estimates of Turnbull et al. (2009) for North American mid-latitudes and set this correction to -0.2 (+/-0.1) ppm (thus resulting in a positive offset) from October to March and to -0.5 (+/-0.3) ppm from April to September. Since the correction term in Eq. 3 is subtracted from the first term, the impact of heterotrophic respiration is to raise estimates of CO_2 ff in both seasons.

The influence of additional sources on Δ^{14} obs is globally variable and has potential contributions from stratospheric intrusion of cosmogenically produced and bomb-era ¹⁴C (e.g. Levin et al., 2010; Graven et al., 2012a), nuclear reactors (e.g. Graven and Gruber, 2011), biomass burning (e.g. Schuur et al., 2003; Vay et al., 2011), and the oceanic-atmosphere disequilibrium (e.g. Sweeney et al., 2007; Muller et al., 2008). However, model-based estimates of the Δ^{14} C signal (not including those from nuclear emissions) in the conterminous United States (Miller et al., 2012) show that these terms contribute very little relative to the spatial gradients arising from fossil fuel combustion. Graven and Gruber (2011) argue that in the eastern United States nuclear contributions may be significant, but they predict near-zero nuclear influence in most of the western United States, including Colorado. Any contribution from stratosphere or ocean sources at BAO is likely to simultaneously impact the NWR background site and, thus, can be ignored in this analysis. At least one sample was influenced by a biomass burning event, identified by an anomalously high CO/CO₂ff ratio, as well as multiple news reports of poor air quality on that particular day resulting from the Station Fire in southern California in August 2009 (e.g. Brennan, 2009). This sample, along with one other that exhibits an abnormally high CO/CO₂ff ratio is omitted from this analysis. The sample influenced by the wildfire plume was collected Sep 1 2009; the other sample, collected Jan 30 2010, is unusual in that the estimated CO₂bio mole fraction was very large (15 ppm), and about twice the estimated CO₂ff for this sample. The large CO₂bio relative to other samples in the data set suggests the possibility of an undetected stratospheric or biomass burning influence or an unusually large heterotrophic respiration signal, therefore we exclude this point from our analysis. In addition to CO, a large number of other anthropogenic tracers were elevated in this particular sample.

2.6 Bottom-up fossil fuel CO₂ emissions estimates

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To derive top-down emissions estimates for the observed trace gases via tracer/CO₂ff enhancement ratios, we use both county-level and gridded bottom-up fossil fuel CO₂ emissions estimates from the Vulcan data product (v2.2) (Gurney et al., 2009) as a quantitative reference. Vulcan (http://vulcan.project.asu.edu) is a high resolution data product that utilizes a combination of energy, air quality, census, traffic, and digital road statistics to quantify fossil fuel CO₂ emissions for the United States. Until recently, the Vulcan inventory was available only for 2002, but is now updated to include

annual emissions at the county and state level for all years between 1999 and 2008. The gridded high resolution product is currently available only for 2002, however. Since we make use of both the county-level and gridded inventories in our analysis, the Vulcan02 data product is used as the base year for consistency. For the Vulcan02 product, country-wide, emissions are in agreement with the United States Energy Information Administration (EIA) estimates to about 2% even though these emissions were compiled using two different independent statistical data sets (Gurney et al., 2011). At the county level, the estimated uncertainty (1σ) on annual CO₂ff emissions from the VulcanO2 data product is variable, but no more than ~20% (and typically less than ~10%) for any given county (Gurney et al., 2011). To apply the Vulcan02 data product to our analysis period (2009-2010), the Vulcan02 emissions are scaled up to the observation period using the state-level EIA inventory (EIA, 2012), which is currently available through 2009. We use the county-level Vulcan data product for 2003-2008 to constrain the uncertainty in our scaling factor derived from the state-level EIA data. A more detailed description of the scaling procedure and associated uncertainty is provided below (Sec. 3.3).

Vulcan emission rates for CO_2 are given in Table 1 for two source regions that correspond to the N/E and S wind sectors, as defined above (Sec. 2.6). For simplicity we define the N/E wind sector as being influenced primarily by emissions from Weld and Larimer Counties and the S wind sector as being influenced primarily by emissions from Adams, Broomfield, Arapahoe, Jefferson, and Denver Counties (collectively referred to here as the Denver metro counties). Total CO_2 ff emissions, according to VulcanO2, are estimated to be 2.94 Tg C and 7.27 Tg C for the N/E (Weld and Larimer Counties) and S (Denver metro counties) wind sectors, respectively. In Sec. 3.3.1, we consider the uncertainties associated with these assumptions about the geographic area influencing emissions in the two wind sectors.

2.7 Bottom-up trace gas emissions estimates

We compare our top-down emission estimates with bottom-up data, where available, for each tracer species, including: CO (NEI 2008), acetylene (NEI 2005), benzene (NEI 2008 and CFRPS), CH₄ (EDGAR v4.2 2008 and CFRPS), and the C_3 - C_5 alkanes (NEI 2005 and CFRPS). Emissions of C_2 H₂ and the C_3 - C_5 alkanes are estimated from a gridded NEI 2005 inventory of total VOC emissions in combination with the EPA SPECIATE(v4.3) model (EPA, 2011). Additional estimates of certain gases are provided in the CFRPS, which derived a bottom-up inventory for benzene, CH₄, and the C_3 - C_5 alkanes representing emissions from flashing of condensate storage tanks and venting of natural gas wells in Weld County in 2008. These emissions estimates were derived from total VOC emissions estimates from the WRAP/Environ inventory (Bar-llan et al., 2008a, b) in combination with a series of speciation profiles of raw natural gas (for venting emissions) and condensate tank mixtures (for flashing emissions) in the region. From the various speciation profiles available, average emissions were calculated for each along with the maximum and minimum values, thus giving a range of bottom-up estimates (Pétron et al., 2012). As we will discuss below, the CFRPS bottom-up inventory is a compilation of emissions from oil and gas related activities only, but other important sources exist for some species, in particular for benzene and methane.

Table 1 summarizes the bottom-up emission estimates, including the base-year for each inventory. Scaling factors (α) for the trace gases that relate the inventory base-year to the observation period are estimated from population statistics or additional factors. Scaling of the tracer inventories related primarily to mobile emissions (i.e. CO, benzene, and C₂H₂) is calculated in proportion to the rate of increase in population according to statistics from the U.S. Census Bureau. For these species, uncertainty limits for the scaled emissions are assigned as the base year estimates (i.e. no change in emissions) on the low end to an estimate using a scaling factor that is 3 times the population increase on the high end. The one exception to this is for the uncertainty limits for CO emissions. There is evidence that on road mobile CO emissions have decreased in many urban regions over the past 15-20 years despite large population increases, and in Denver, specifically, the CO-to-fuel burnt ratio was observed to have decreased at a rate of about 7% per year between 1999 and 2007 (Bishop and Stedman, 2008). Therefore, the bottom-up CO emissions uncertainty is bracketed at the low end by an emission rate corresponding to a decrease in emissions of 10.5% from 2008 (the inventory base year) to the observation period. For CH₄ and the C₃-C₅ alkanes, emissions in Weld/Larimer counties are scaled by increases in gas and oil production (by total dollar amount), according to statistics from the Colorado Oil and Gas Conservation Commission (COGCC, 2011). Uncertainties for the CFRPS base-year estimates represent the minimum (un-scaled) and maximum (scaled) values, given a set of 3 speciation profiles based on measured raw natural gas composition data for 77 wells (COGCC, 2007) and a set of 16 modeled flash emissions profiles for condensate tanks (provided by CDPHE, see details in Pétron et al., 2012). These uncertainties are modified here to reflect the additional uncertainty resulting from upscaling to the observation period. Since virtually all the gas and oil wells in the Weld/Larimer county region are located in Weld county (99.8% of the total production, according to the COGCC), the CFRPS estimates, which correspond to Weld county emissions alone, are taken to represent the total Weld/Larimer emission rate. For the South wind sector, where the oil and gas industry is likely the source of only a small percentage of emissions, the EDGARv4.2 (EDGAR hereafter) 2008 CH₄ and NEI 2005 C_3 - C_5 inventory emission estimates are scaled in accordance with population increases, with uncertainties assigned as described above for trace gases related to mobile emissions. We acknowledge that scaling up of these trace gas estimates is an unconstrained approximation, especially for species such as benzene and methane, which have multiple unrelated sources that likely do not scale linearly. For this reason we have taken a conservative approach in assigning scaling uncertainties. It is important to note, however, that the inventory base year estimate is always within the uncertainty brackets of the scaled inventory values, thus allowing the reader to evaluate the top-down and bottom-up comparison independent of any scaling assumptions made here.

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3. Results and Discussion

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3.1. Δ^{14} C and CO₂ff Time Series

The results of the ¹⁴CO₂ analyses are shown in Fig. 2a with values ranging from -19.4 to 50.5‰. The time series runs from June 2009 to September 2010, overlapping with the observation period of the CFRPS,

413 where observations (from the same set of flask samples) up through the spring of 2010 were included their top-down emission calculations. Excursions of $\Delta^{14}CO_2$ at BAO (relative to the NWR background site) 414 towards lower values signify the addition of recently emitted fossil fuel CO2 to the sampled air mass. As 415 416 described in Sec. 2.5, the CO₂ff mole fraction can be quantified using Eq. 3, with an uncertainty of 1.2 ppm based on propagation of the analytical uncertainty in Δ^{14} CO₂ for both Δ^{14} obs and Δ^{14} bkg (the 417 uncertainty in CO_2 terms is small relative to those for $\Delta^{14}C$). Performing this calculation for each BAO 418 419 observation in Fig. 2a gives CO₂ff mole fractions that range from below the 1.2 ppm detection limit up to 420 25 ppm. There are instances of negative CO₂ff values (14% of all samples), which is not physically 421 realistic, in the dataset. All but 5 of these samples (3% of the entire data set) lie within the 1σ envelope 422 around zero and only 1 sample (-3.3 ppm) lies outside of 2σ .

The most obvious feature of the CO_2 ff variability is that mole fractions are high and variable in the winter months and relatively constant and lower, on average, during the summer months (Fig. 2b). This trend is qualitatively consistent with shallow, and variable, boundary layer heights in the winter and deep boundary layers in the summer. Boundary layer height is driven by a number of complex meteorological and topographical variables, but largely by surface sensible heat flux, which is of course much lower during the winter. Tracer/tracer ratios are expected to be much less sensitive to variability in boundary layer height since the dilution and mixing of co-located emissions will impact the different tracers equally. As we describe below, observations of a set of tracer/ CO_2 ff ratios are consistent with this expectation.

3.2. Variability in tracer/CO₂ff enhancement ratios

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When sources of trace gas emissions are co-located with fossil fuel combustion sources, an analysis of the trace gas abundances relative to CO₂ff provides a means to better understand the variability in the mix of emission sources influencing the site independent of the dilution and mixing dynamics that impact absolute mole fractions. An alternate explanation could be that emissions of all the trace gases have similar seasonal cycles to CO₂ff. Tracer/CO₂ff enhancement ratios are calculated here by taking the median of individual tracer/CO₂ff ratios after subtracting the background from each trace gas. The median emission ratio derived from all samples provides a more robust estimate of the apparent tracer/CO₂ff ratios than that determined from either a linear regression slope or an arithmetic mean, which give estimates that are overly sensitive to ratio outliers that can result from signals due to air masses in which emissions of various sources are not well mixed (Miller et al., 2012). Samples are only used in the ratio calculation when estimated CO₂ff is above the 1.2 ppm 1σ detection limit. Uncertainties in the median ratios are 95% confidence intervals, defined as the 2.5-97.5 percentile range (~2σ confidence) from a distribution of 500 estimates of the median from a randomized re-sampling of the data (boot-strapping with replacement). We also estimated the uncertainty in the tracer/CO₂ff enhancement ratios associated with measurement uncertainty (both for the trace gas and CO₂ff) and found that these uncertainties (at 2σ) were comparable or lower than the boot-strap approach in all cases. A measure of the appropriateness of this approach is estimated by calculating the coefficient of determination (r²) from a linear regression of tracers vs. CO2ff; a high r² suggests that emissions of the tracers are appreciably co-located with fossil fuel combustion sources. Results from the tracer/CO₂ff enhancement ratio calculations (with associated uncertainties and r² values) are detailed in Table 1.

Background observations for the different trace gases are taken from one of two nearby sites in the NOAA-GMD global network, either NWR (CO and CH_4) or from flights at CAR (acetylene, benzene, and the C_3 - C_5 alkanes). CO and CH_4 observations are available from both sites and we confirmed that the enhancement ratio estimates are not appreciably sensitive to the selection of background site (differences between 7% and 15% in derived enhancement ratios).

The sensitivity of this analysis to the prescribed heterotrophic respiration correction to CO_2 ff (Eq. 3) was determined by recalculating the tracer/ CO_2 ff ratios with this correction term doubled, in one case, and set to zero in another. The ratios estimated from this sensitivity test were within the 95% confidence intervals in all but one case. The lone exception was the CH_4/CO_2 ff ratio for the N/E wind sector, where the recalculated ratio using a respiration correction of zero is 34.1 ppb/ppm compared to the base case upper confidence limit of 33.4 ppb/ppm, extending the upper uncertainty limit by only 2%. Thus we consider the uncertainty in the heterotrophic respiration correction to CO_2 ff a largely insignificant source of error in our analysis. Given the relative lack of vegetation in the region surrounding BAO, it is more likely that the prescribed respiration correction is biased high rather than low, which would result in CO_2 ff values that are biased high and tracer/ CO_2 ff enhancement ratios that are biased low.

While variability in the absolute mole fractions of the tracers and CO₂ff has a strong seasonal dependence (e.g. Fig. 2b), with larger enhancements observed in the winter than the summer, there is no apparent seasonality to any of the considered tracer/CO₂ff enhancement ratios, suggesting that boundary layer dynamics are largely what are driving the seasonality in measured atmospheric mole fractions. As we discuss below, the primary source of variability in the tracer/CO₂ff enhancement ratios at BAO is due to changes in wind direction, with significantly different tracer signatures observed when air is transported from Weld and Larimer counties to the north and east (N/E) vs. from the Denver metro counties to the south (S). A relatively small number of samples in this data set arrived from the W wind sector, most of which did not have very strong enhancements of CO₂ff or other trace gases. We therefore ignore these samples in our analysis.

Previous work (Pétron et al., 2012) found significant differences in mole fractions of alkanes, including CH_4 , C_3H_8 , n- C_4H_{10} , i- C_5H_{12} , and n- C_5H_{12} , observed at BAO as a function of wind direction and related differences in emission sources. Benzene was also found to be enhanced in air masses arriving from the N/E. These differences were attributed to oil and gas production in Weld County, to the northeast of BAO. As shown on the map in Fig. 1, the majority of these wells are located in Weld County (SkyTruth, 2008), from which 17.9 million barrels of oil and 5.7 billion cubic meters of natural gas were produced in 2009 (COGCC, 2011). Other potential sources of CH_4 include cattle feedlots, landfills, waste water treatment plants, and natural gas processing plants. The transportation or mobile sector contributes significantly to a subset of the gases considered: CO, C_2H_2 , C_6H_6 , and i- C_5H_{12} (Watson et al., 2001). This sector likely contributes significantly to emissions from the Denver metro counties, but there are also significant mobile emissions in the N/E wind sector from Interstate 25, the main north-south route in Colorado, as well as in a number of population centers, including Fort Collins, all located due north of BAO, in Larimer County. Table 2 summarizes the expected sources of the trace gases evaluated in this analysis, along with their expected atmospheric lifetime with respect to oxidation by OH. Lifetimes of the tracers considered range from 3 days (pentanes) to 7 years (CH_4) (at a modest OH molecular density

- of 10⁶ cm⁻³). The oxidation of tracers can potentially reduce the observed enhancement ratio resulting in
- a lowering of the presumed emission ratio; however, as we will discuss, we find no evidence of sufficient
- 495 OH chemistry to impact the tracer/CO₂ff ratios discussed here.

3.2.1. Carbon Monoxide

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- 497 Fig. 3a shows the relationship between CO and CO₂ff for each sample. Fits of a linear regression are
- 498 included in the Fig. 3a for the N/E and S wind sectors, giving r^2 values of 0.75 (n = 44) and 0.85 (n = 25),
- 499 respectively. As detailed in Table 1, and shown in Fig. 4, the point-by-point analysis of these
- observations show median (with 2σ confidence intervals) CO/CO₂ff enhancement ratios of 9.3 (8.1 –
- 501 10.6) and 10.0 (6.7 13.3) for the N/E and S wind directions, respectively. For all wind sectors combined,
- 502 the median ratio is 9.3 (8.3 10.8) ($r^2 = 0.81$).
- The observed ratios from both wind sectors are similar to the values of 6.8 ± 2.2 and 11.7 ± 5.5 ppb/ppm
- 504 calculated at Niwot Ridge from two samples originating from the Denver area via upslope winds in 2004
- 505 (Turnbull et al., 2006). Our estimates are somewhat lower, however, than previous reported values of
- 506 CO/CO₂ff in Denver, where ratios were derived from linear correlations across 4 different aircraft flights
- 507 (~4-6 samples per flight) in May and July of 2004 (Graven et al., 2009). The observed ratios from these
- 508 flights ranged from 14 27 ppb/ppm. While the well documented (e.g. Bishop and Stedman, 2008)
- reductions in CO emissions from mobile sources between 2004 and 2009 (part of a much longer term
- trend across most of the country) could be a factor in the lower enhancement ratios observed here, the
- long term data set from BAO provides a more robust estimate of the CO/CO₂ff ratio than either of these
- short-term studies where small errors in individual data points could result in a large difference in the
- estimated ratio and where short term variability could have a strong influence. For comparison with
- these short term data sets, observed ratios of CO/CO₂ff for individual samples from the south wind
- sector at BAO range from 3.6 to 13.5 ppb/ppm (1 σ), with a maximum observed value of 20 ppb/ppm
- 516 (not including the sample impacted by biomass burning). Differences in the influencing area of emissions
- between the two studies may also play a role in the observed differences.
- The main anthropogenic sources of CO in Colorado, and in much of the US, are from on-road gasoline
- vehicles in the mobile sector (66%) and from non-road gasoline-based equipment (26%) (NEI, 2008).
- 520 While the on-road and non-road sectors account for 92% of total CO emissions in Colorado, these
- 521 sectors contribute only 29% of the total CO₂ff emissions according to the Vulcan08 data product (Gurney
- 522 et al., 2009), suggesting that the CO/CO₂ff emission ratio from other combustion sources is very small.
- 523 Similar CO/CO₂ff ratios for both N/E and S wind sectors, therefore, suggests a similar contribution of on-
- road and non-road CO₂ff sources in both Weld/Larimer counties and the Denver metro counties,
- 525 consistent with the Vulcan emissions which estimates that the on-road plus non-road sectors, the
- dominant CO contributors, combine for 29% and 41% of the total CO₂ emissions, for Weld/Larimer and
- 527 Denver metro respectively (Gurney et al., 2009). This is in contrast to CH₄ and other trace gases, as we
- 528 discuss below, where there is a clear enhancement due to non-combustion sources related to oil and
- 529 gas production.

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3.2.2. Methane

531 We find significant differences in the mole fraction of CH_4 relative to CO_2 ff depending on wind direction. This can be seen in the correlation plot of CH₄ with CO₂ff (Fig. 3b), where filtering by wind sector results 532 in two highly correlated relationships (r² of 0.87 and 0.69 for Weld county and the metro Denver 533 counties, respectively); however, the slope is higher by a factor of 3 in the N/E wind sector relative to 534 535 the S wind sector. The implication of this is that emissions of CH₄, relative to CO₂ff, are 3 times higher in 536 the N/E sector than the S sector. The added source of methane influencing air samples arriving from the 537 N/E are likely due to fugitive emissions of raw gas and flashing emissions from condensate storage tanks 538 (Pétron et al., 2012). Most of the operations are located in Weld County in the DJB. Condensate tanks 539 store a semi-liquid mix of hydrocarbons separated from raw natural gas; flashing emissions occur when 540 condensate liquids experience a drop in pressure, causing entrained gas to escape. Venting/fugitive 541 emissions can originate at the wellhead and occur when new gas or oil wells are drilled and completed 542 or when existing wells are vented or repaired. Additional sources of methane from the oil and gas sector 543 may include pipeline leaks, pneumatic devices and pumps, and incomplete combustion in compressor 544 engines. Cattle feedlots, wastewater treatment plants, and landfills are also present in the region and 545 likely contribute significantly to the CH₄ signal at BAO.

While venting and flashing are not associated with combustion, except insofar as they are co-located with heavy equipment and compressor engines, there is the possibility of entrained CO_2 ff being coemitted from natural gas wells. CO_2 can be a small fraction (3-5% by mass) of raw natural gas (COGCC, 2011), but constitutes only a negligible fraction (<0.1%) of total Weld/Larimer county CO_2 emissions, based on the estimates of Pétron et al. (2012). This suggests that the correlation of CO_2 ff with CH_4 is due primarily to similarly located, but separate processes. Further evidence of this can be found in a consideration of multiple tracer/ CO_2 ff ratios, as we will discuss below.

3.2.3. Other trace gases

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To further understand the differences in emissions between the two wind sectors, we consider the differences across a series of tracer/CO₂ff ratios. Fig. 5 shows the difference in median tracer/CO₂ff ratios for CO, C₂H₂, CH₄, C₃-C₅ alkanes, and benzene when winds are from the N/E and from the S. Like CO, C₂H₂ is emitted overwhelmingly from combustion sources, while the other trace gases are emitted either from non-combustion sources (C_3H_8 , n- C_4H_{10} , and n- C_5H_{12}) or from a combination of sources. Both CO and C₂H₂ show no appreciable dependence on wind direction, consistent with the idea that both gases are emitted primarily from combustion processes that are common to Weld/Larimer counties and the Denver metro counties. The median enhancement ratio of C₂H₂ to CO₂ff observed at BAO (N/E and S combined) is 43.7 (38.3-55.4) ppt/ppm (16^{th} -84th percentile range) (r^2 = 0.81), which is consistent with observations from two previous studies in different locations: 52 (45-59) ppt/ppm downwind of Sacramento, CA (Turnbull et al., 2011) and 45.9 (28.6 – 102.9) ppt/ppm off the east coast of the United States during winter (Miller et al., 2012). This consistency suggests a relative insensitivity of this ratio to a particular mix of emission type across the United States, an important criterion if one were to consider using C_2H_2 as a proxy for CO_2ff in the absence of $\Delta^{14}CO_2$ observations. However, the large spread observed in the enhancement ratio off the eastern U.S. coast by Miller et al. (2012) (as reflected by the 16th and 84th percentiles of the distribution of observed ratios) suggests that there can be more variability in this ratio than indicated by the range of median values alone. Additional research is

required to better evaluate the potential for using C_2H_2 as a secondary CO_2 ff tracer and whether it would prove advantageous over the use of CO (Turnbull et al., 2006; Levin and Karstens, 2007), which may be problematic in locations where significant is situ production results from VOC oxidation.

As with CH_4 , there are significant differences in the tracer/ CO_2 ff enhancement ratios for the C_3 - C_5 alkanes and benzene with wind direction, which suggests that enhanced emissions of these chemicals are associated with gas and oil operations (Bar-Ilan et al., 2008a, b; Pétron et al., 2012). In general, ratios of C_3 - C_5 alkanes are enhanced relative to CO_2 ff by about a factor of 10 in the N/E wind sector compared to the S wind sector. Benzene is enhanced in the N/E wind sector compared to the S wind sector by a factor of 1.6. Despite the significant non-combustion sources of the VOCs related to gas and oil production, we see very good correlations of these species with CO_2 ff in air arriving from the N/E ($r^2 > 0.75$), an indication of integration of emissions by air mass mixing or substantial co-location of combustion sources with gas and oil wells and condensate tanks.

3.3. Estimating emission magnitudes

From the observations described above as well as those reported in the CFRPS, it is clear that air sampled at the BAO tall tower is influenced by emissions on local and regional scales. Changes in wind direction at the site result in these local emissions coming from one of two primary source regions: (1) gas and oil operations to the north and east and (2) the Denver metro region to the south. Given the distinct geographical separation of sources, we use the wind sector specific observations, in conjunction with county-level CO_2 emissions from the Vulcan data product (Gurney et al., 2009) as a means to constrain emissions for these trace gases using a top-down approach.

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$$E_x = E_{CO2ff}(1 + \alpha/100)R$$
 Eq. 4

Eq. 4 describes the annual average top-down emissions for a series of trace gases (E_x), where R is the median observed tracer/ CO_2 ff ratio, E_{CO2ff} is the annual average Vulcan CO_2 ff emission rate for the region of interest, and α is a scaling factor that is designed to account for changes in emissions from the emission base year to the observation period. For CO_2 ff emissions, this factor is equal to the change in emissions (expressed as a %) for the EIA inventory for Colorado state between 2002 (the Vulcan base year) and the most current EIA inventory year, 2009. Equation (4) is applied independently to the N/E and S wind sectors for each tracer, with R calculated for the N/E and S wind sectors paired with E_{CO2ff} estimates for Weld/Larimer counties and the metro Denver counties, respectively. Since α is based on state wide changes in E_{CO2ff} , this scaling factor is equivalent for both wind sectors. Tracer/ CO_2 ff ratios (R) are calculated as discussed in Sec 3.2. Table 1 summarizes the parameters used to calculate E_x for Weld/Larimer counties and the Denver metro counties. Note that the observation period spans two summers (and one winter) and thus any seasonal bias in the observed value R would lead to summer emissions being over-represented in the estimates of E_x . From the available data, however, we can detect no significant differences (with respect to the 2σ confidence intervals) in R with season for any of the trace gases considered.

Uncertainties in R (as described in Sec 3.2), E_{CO2ff} , and α are considered (among other factors) in the estimation of top-down emissions. The scaling term, α , is 2.8% for the state of Colorado according to the

EIA inventory. While this scaling term indicates almost no change between 2002 and 2009 emissions, in actuality, emissions increased by 9% by 2007 and then decreased over the next 2 years (presumably related to the economic downturn in the United States during this period). Similar trends are observed in the county level Vulcan emissions over this time period, though the peak year in both Denver Metro and Weld/Larimer counties occurs earlier than 2007. Using changes in the annual EIA-based Colorado emissions to scale the Denver Metro and Weld/Larimer Vulcan02 estimates, gives, in general, very good agreement with the Vulcan estimates for these counties from 2003-2008 (to within 10% for any given year and about 5% on average). The Vulcan02 uncertainties (1 σ) for the individual counties considered here are of similar order, ranging from 4.6% to 10.6% (K. Gurney, personal communication), with less uncertainty associated with the combined larger county "sectors" that we use in our wind sector analysis. Doubling these uncertainties (to be consistent with our 2 σ analysis) for the two wind sectors results in differences from the central estimate of 7% (upper estimate) and 11% (lower estimate) for Weld/Larimer counties and 7% (upper) and 10% (lower) for the Denver metro counties. We therefore assign a conservative uncertainty of $\pm 20\%$ to the scaled bottom-up CO₂ff emissions estimates in this analysis, folding in uncertainty in both E_{CO2ff} and σ .

The calculated top-down estimates (E_x) are given as a central estimate or 'best guess' for the annual emissions plus 95% confidence intervals calculated by propagation of the uncertainties described above. For species having sources that are not necessarily spatially correlated with emissions of CO_2 ff (e.g. venting and flashing emissions), it is possible that the top-down emissions estimates are biased in some way. However, it is not possible to predict what direction this bias would be in since it is equally likely for an enhanced CO_2 ff sample to arrive at the site with no accompanying trace gas enhancement as it is for the trace gas to be enhanced but not CO_2 ff. The boot-strap determination of uncertainties for the enhancement ratio puts a reasonable constraint on the impact of less than perfect correlations between the trace gases and CO_2 ff. Figures 6(a) and 6(b) summarize the top-down estimates and confidence intervals (whiskers), along with the available bottom-up estimates (from multiple sources) and top-down estimates from the CFRPS, for each trace gas species considered.

3.3.1 Spatial Considerations

Additional uncertainty in E_{CO2ff} arises as a result of our simplistic assumptions regarding the geographic footprint (area of emissions) influencing the observations. Obviously, the emissions influencing the observations are not strictly confined to the county boundaries that we have selected, based on the simple wind sector analysis. This matters only to the extent that the spatial distribution of tracer/ CO_2ff emission ratios varies between the presumed footprint and the actual footprint. This may be an issue especially for emissions estimates in the N/E wind sector where VOC and CH_4 emissions from the DJB are primarily confined to within Weld County while CO_2 emissions are likely significant over a larger spatial scale. For example, there are significant CO_2 emissions along the I-25 corridor (in Larimer County) to the north of BAO, where there are relatively few active gas wells (see Fig. 1). Whereas CO_2 emissions are significant in both Weld (55%) and Larimer (45%) counties (according to Vulcan2008), the vast majority (~99.8%) of natural gas and oil production (and associated emissions, presumably) in the two counties is confined to Weld County (COGCC, 2011). Thus, the top-down emissions estimates of the trace gases

from oil and gas production will be sensitive to our assumptions about the geospatial scale of the observations, specifically, whether the observations are influenced by emission fluxes only in Weld County or across a larger area that extends into the eastern part of Larimer County or other locations where CO_2 ff emissions are significant.

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An available constraint on these assumptions is the top-down estimates of CH₄ from the CFRPS (Fig. 6a), where bottom-up estimates of C₃H₈ and CH₄ emissions from condensate tanks were used as a quantitative reference. Since C₃H₈ and CH₄ are strongly correlated at BAO and are emitted from the same processes related to the oil and gas industry in the region, and thus distributed similarly in space, the CFRPS top-down estimates are somewhat less sensitive to assumptions about the spatial extent of the observation footprint, even though some uncorrelated sources of CH₄ from cattle feedlots, for example, may be significant. Using the CFRPS top-down CH₄ estimate of 143 (72 – 271) Gg yr⁻¹ (which has been scaled up to 2009-2010) and our observed CH₄/CO₂ff ratio of 30 ppb/ppm, we calculate corresponding CO_2 ff emissions of 3.6 (1.8 – 6.8) Tg C yr⁻¹. The central estimate of this calculation is about 20% greater than the Larimer plus Weld county CO₂ff emissions from the scaled Vulcan data product. For Weld County alone, the central estimate (1.7 Tg C yr⁻¹) is just below the lower end of the range of estimates. The range of top-down CH₄ estimates from L12 is considerably lower than the CFRPS estimates, resulting in inferred CO_2 ff emissions of 1.1 – 2.2 Tg C yr⁻¹ (using "realistic errors") or 1.1 – 3.3 Tg C yr⁻¹ (using "conservative errors"). The inconsistencies between the CFRPS and L12 are significant, and it is beyond the scope of this study to pass judgment on the validity of one estimate over the other, so we simply expand the range of top-down CO₂ff estimates to be consistent with both studies.

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The full range of CO₂ff emissions estimates (1.1-6.8 Tg C), as derived from the CFRPS and L12 top-down CH₄ estimates, is much larger than the 20% uncertainty that we prescribe for the county-level bottomup emissions, and thus translates to a large uncertainty in the spatial extent of emissions influencing observations at BAO. We expect that our analysis of trace gas emissions related to the oil and gas industry in the N/E wind sector to be particularly sensitive to the uncertainty in the spatial extent of observations, due to the relatively poor spatial correlation between CO₂ff sources and oil and gas operations across the whole of Weld and Larimer counties. Thus, we use this range of E_{CO2ff} estimates (1.1 - 6.8 Tg C) to set the uncertainty limits on our determination of emissions for the trace gases related specifically to the oil and gas industry in Weld County (CH₄, benzene, and the C_3 - C_5 alkanes). These upper and lower bounds should be interpreted as representing an upper limit of the uncertainty resulting from having no independent information about the spatial extent of this analysis. Therefore, while this range is large and does not provide an independent estimate of emissions of these trace gases (due to the reliance on CFRPS and L12 calculations), it does allow us to set a conservative range of geospatial scales over which a series bottom-up vs top-down comparisons can be made. We will revisit this discussion in Sec 3.3.4 by performing a sensitivity test of our top-down analysis to different assumptions about the spatial extent of our observations, using CH₄ as an example. For CO and C₂H₂ and for all emissions estimates for the Denver metro counties we assume that tracer emissions are more spatially correlated with CO₂ff and assign uncertainty as described in Sec. 3.3.

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3.3.2 Carbon Monoxide

We estimate annual CO emissions to be 65.8 (50.1 - 81.7) Gg yr⁻¹ CO for Weld/Larimer counties and

175.1 (106.8 - 241.2) Gg yr⁻¹CO for the metro Denver counties. The NEI08 estimates for these regions

are 120.1 Gg CO and 362.1 Gg yr⁻¹ CO, corresponding to overestimates by a factor of 1.8 and 2.1, for

694 Weld/Larimer and metro Denver counties, respectively (Fig. 6). Total emissions for the two regions are

overestimated by a factor of 2.1 in the NEI bottom-up inventory; the range of uncertainty in the

696 combined top-down estimate corresponds to a range of 1.7 – 2.8 for the overestimate of the NEI

697 bottom-up inventory. These values are consistent with prior studies evaluating the accuracy of the NEI

698 CO emissions (Parrish, 2006; Hudman et al., 2008; Miller et al., 2008; Turnbull et al., 2011; Miller et al.,

699 2012).

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700 Comparing the results of different studies where radiocarbon observations were used to derive

CO/CO₂ff ratios provides some insight into the ubiquity of the overestimate of CO emissions in the NEI

inventory. A survey of observed CO/CO₂ff ratios from different locations, including Sacramento, CA

703 (Turnbull et al., 2011), Denver (Turnbull et al., 2006; Graven et al., 2009), Irvine, CA (Djuricin et al.,

704 2010), and off the eastern coast of the United States (Miller et al., 2012), reveals regional differences in

the agreement between observations and bottom-up estimates (Fig. 4). The observations off of the

706 eastern seaboard of the United States point to an overestimate of NEI CO emissions (Miller et al., 2012),

707 consistent with the BAO observations and with the numerous literature examples finding CO emissions

are over-estimated across the US (Hudman et al., 2004; Parrish, 2006). However, both California-based

studies find that the observed CO/CO₂ff ratio closely matches the estimates from California bottom-up

710 inventories (Fig. 4). The Sacramento results were compared with bottom-up estimates from both the

711 California Air Resources Board and NEI05 inventories by Turnbull et al. (2011). The authors of that study

found good agreement with the CARB 2008 inventory and a factor of about two overestimate in the

NEIO5 inventory. From our analysis, it appears as though this overestimate in the NEI inventory for

California has been corrected in the 2008 release, perhaps a result of adopting the CARB estimates, as

previously suggested by Turnbull et al. (2011). Similarly, the observations in Irvine (Djuricin et al., 2010)

are in good agreement with the NEIO8 inventory for the LA Basin (Los Angeles, Orange County, San

717 Bernardino, and Riverside counties). From the BAO observations and those off the east coast of the US,

it appears as though this correction was not made for the rest of the country.

719 As with the previous Denver observations, the Sacramento and Irvine observations are representative of

shorter time periods: the Sacramento observations were compiled from linear correlations of several

721 samples collected during 2 aircraft flights, while the Irvine observations were from 3 discrete samples

722 collected over a few different months at a surface site on the campus of UC Irvine. The Miller et al. study

provides a longer term average, similar to the BAO observations, but is more representative of northeast

US regional-scale ($^{10^5}$ - $^{10^6}$ km²) sources rather than the local-to-regional ($^{10^3}$ - $^{10^5}$ km²) influence

at BAO. Given the large differences in scale relevant to each of these studies, the comparisons of

726 CO/CO₂ff ratios in the various locations are not necessarily conclusive. However, as we will discuss in Sec

4.1, we find that a closer inspection of the NEIO8 inventory reveals a significant difference in the CO

inventory in California vs Colorado that is not supported by observations.

729 In situ production or loss of CO could potentially bias these results. The most likely scenario would be

the production of CO from the oxidation of VOCs by OH (Griffin et al., 2007) and can be significant in

- 731 some locations, especially during summer when oxidation rates are intensified and biogenic VOC
- 732 emissions are high. At BAO, we do not see an appreciable difference in the CO/CO₂ff ratio from winter to
- summer, which suggests a minimal influence of photochemistry on CO abundance at BAO. The
- atmospheric lifetime of CO is sufficiently long (~50 days) that its impact is negligible to this analysis.

3.3.3 Acetylene

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- Acetylene emissions are estimated to be 0.28 (0.21 0.40) Gg yr⁻¹ in Weld/Larimer counties and 0.78
- 737 (0.54 1.0) Gg yr⁻¹ in the Denver Metro counties. These values are higher than the bottom-up estimates
- by factors of 1.5 (1.1-2.1) and 1.4 (0.9-1.7) for Weld/Larimer and Denver metro, respectively (Fig. 6). In
- 739 contrast to carbon monoxide, there has been very little evaluation of C₂H₂ emissions inventories in the
- 740 United States. Warneke et al. (2007) compared the C₂H₂:CO ratio from observations in Boston, New
- 741 York, and Los Angeles to that in the NEI99 emissions database, and found the ratio to be
- underestimated in each location, suggesting a systematic underestimation of acetylene emissions by the
- 743 NEI database. However, it is unclear whether this underestimation of the C₂H₂:CO ratio is a result of an
- underestimate of C_2H_2 or an overestimate of CO (as detailed above). With observations of $^{14}CO_2$, the
- 745 C₂H₂ inventory can be directly evaluated independently of any biases in the CO emissions inventory.
- There have been two recent examples comparing top-down estimates of C₂H₂ emissions in the United
- States to bottom-up inventories using ¹⁴CO₂ observations: Miller et al. (2012) estimated C₂H₂ emissions
- 748 for the entire United States (assuming northeast ratios were nationally representative) and found
- relative agreement (within 6%) with the NEI05 C₂H₂ emissions inventory (the same gridded inventory
- used for comparison in this study and described above); Turnbull et al. (2011) published a comparison of
- 751 C₂H₂:CO₂ff ratios from observations of the Sacramento urban plume with that calculated from bottom-
- up inventories and found a ~30% underestimate of C₂H₂ in the NEI 2005 inventory for Sacramento, CA.
- Additional ¹⁴CO₂ observations co-located with C₂H₂ in more locations and comparison with
- 754 contemporaneous NEI values are required to come to any definitive conclusions regarding the accuracy
- of C₂H₂ emissions in the NEI database. The use of C₂H₂ as a secondary fossil fuel tracer or a proxy for
- 756 CO_2 ff seems promising, however, and given the limited attention that evaluation of C_2H_2 emission
- 757 sources have received in the literature, further studies are recommended.

758 **3.3.4 Methane**

- As discussed in Sec. 3.2.2, we sample different sources of CH_4 , as well as benzene and the C_3 - C_5 alkanes,
- 760 relative to CO₂ff emissions with changing wind direction. The oil and gas operations to the northeast of
- 761 BAO, primarily in Weld County, significantly impact concentrations of these species in air at BAO (Pétron
- et al., 2012), consistent with our findings here. Additional sources, including cattle feedlots, waste water
- treatment plants, and landfills are likely also important in this region.
- Annual emissions of CH_4 are estimated to be 121(40-272) Gg yr⁻¹ and 98 (55 138) Gg yr⁻¹, for
- 765 Weld/Larimer counties and the Denver Metro counties, respectively. Bottom-up estimates from the
- 766 CFRPS for the former region are 71 (46-100) Gg yr⁻¹, within the lower uncertainty bracket of our top-
- down estimate (40 Gg yr⁻¹), but considerably below the central top-down estimate (121 Gg yr⁻¹).

As discussed in Sec. 3.3.1, there is significant uncertainty in the spatial extent of emissions influencing observations at BAO, creating ambiguity when defining the geographic constraints for comparing the top-down and bottom-up estimates. We can explore this issue in a more quantitative way by comparing the top-down and bottom-up estimates across a range of geographic areas, using gridded inventories of CO₂ff and CH₄. To do this we convolve an expanding series of hypothetical footprints or "influence functions" onto the gridded Vulcan02 inventory (0.1° x 0.1°), where an influence function is defined as the sensitivity of an observation at a single point in space to fluxes at any given upwind location. For each hypothetical footprint, we can estimate a different value of E_{CO2ff} from which a top-down emission rate can be estimated for CH₄ using Eq (4), with R equal to the observed enhancement ratio. A series of hypothetical footprints is considered that vary in size from 1 x 1 Vulcan grid cells (at 0.1° x 0.1° resolution) to 30 x 30 grid cells and are constructed such that the sensitivity at the receptor to emissions within a given grid-cell decays exponentially with distance from the receptor. Thus the emissions in gridcells close to the tower are always weighted more heavily than those farther away. These hypothetical footprints are not intended to accurately represent atmospheric transport; rather, they are intended to be used to explore the sensitivity of the top-down vs. bottom-up comparison to spatial assumptions across a range of potential influencing regions. Figure 7a shows the weighted CO₂ff emissions across a footprint (13 x 13 grid cells) extending to the north and east that would give approximate agreement with the top-down CH₄ estimate from the CFRPS (143 Gg yr⁻¹). Figure 7b shows the weighting function used to estimate the emissions in Fig 7a. From the full series of hypothetical footprints considered, we find that a range of footprints between 4 x 4 Vulcan grid cells and 26 x 26 grid cells are consistent with the full range of CFRPS and L12 top-down CH₄ estimates (72 – 271 Gg yr⁻¹). This corresponds to an effective area of 2200-74,000 km². An analogous series of footprints are generated for the S wind sector, as well.

The same series of footprints are then convolved with the gridded (0.1° x 0.1°) CH₄ inventory (EDGAR) to provide a bottom-up emission rate for comparison with each top-down estimate. The bottom-up emissions are scaled up by 10% in the N/E wind sector according to changes in oil and gas production between 2008 and 2009-2010 (COGCC, 2011) and by 1.7% for the S wind sector according to population increases in the Denver metro counties. Figure 8 shows the ratio of bottom-up to top-down CH₄ estimates with changes in the effective footprint size for both the S and N/E wind sector. Uncertainties associated with these calculations are carried through from the 95% confidence intervals for the top-down estimates, taking into account uncertainties in the observed tracer/CO₂ff ratio, E_{CO2ff} , and α . To reiterate, these hypothetical footprints are not expected to be an accurate indication of atmospheric transport to the BAO tower, but rather they provide a sense of how sensitive the top-down vs bottom-up comparison is to any presumptions about the geographic region over which emissions are influencing observations at the tower. More directly, this exercise gives an indication of the spatial heterogeneity of CH₄/CO₂ff emission ratios in the bottom up inventory for locations directly upwind of the tower and how this variability influences our conclusions about the accuracy of the bottom-up inventory.

According to the results shown in Fig. 8, the bottom-up/top-down comparison for CH_4 is less sensitive to assumptions about the effective emissions footprint for observations in the S wind sector than in the N/E wind sector. With the exception of only the smallest and largest hypothetical footprints considered,

the bottom-up emissions for the S wind sector are always within 7% of the top-down estimates, and there is general agreement (at >95% confidence) with the EDGAR inventory for urban sources across the entire range of hypothetical footprints considered. A footprint extending to the south and east over an area of 30,000 km² corresponds roughly to the area over which the CO₂ff emissions are equivalent to that specifically from the Denver metro counties. Over this same area, the EDGAR inventory estimates CH₄ emissions of 99 (98-103) Gg yr⁻¹ (with the range of estimates reflecting uncertainties in the scaling parameter, α), in very good agreement with the top-down observations. This bottom-up estimate is included in Table 1 and Figure 6b for completeness.

As discussed above, we estimate that the effective BAO footprint for observations in the N/E wind sector is between 2200 and 74,000 km 2 (or some area of equivalent CO $_2$ emissions). The estimates in Fig. 8b show that in between the limits set by the uncertainty in the effective footprint size, the EDGAR CH $_4$ emissions range from 36%-84% of the top-down estimates from observations in the N/E wind sector. At the high end of this disagreement, the presumed emissions footprint is sufficiently small (4 x 4 grid cells) that uncertainties in both the CO $_2$ ff and CH $_4$ gridded inventories are likely high, and a 64% disagreement is probably not unexpected.

Between 15,000 and 35,000 km², the bottom-up estimate is just within or just outside the lower 95% confidence limit of the top-down estimate. In general, however, the data suggest that emissions of CH₄ are underestimated in the bottom-up inventory. If we remove assumptions about scaling-up the bottom-up inventory from the inventory base year to the observation period, the differences between bottom-up and top-down increase by 10%. One interpretation of this underestimate could be that additional scaling beyond the prescribed 10% is required to account for changes in CH₄ emissions from the inventory base year (2008) to the study period, however we show below that this is likely not the only source of error in the bottom-up inventory. Further, it is likely that emissions have either decreased or scaled non-linearly with production since 2008 due to state regulations requiring the use of low-bleed pneumatic devices (Sgamma, 2012).

At a footprint size (13 x 13 gridcells) that approximates the Weld/Larimer county-scale top-down estimate, the oil and gas sector in the EDGAR inventory contributes about 30 Gg yr⁻¹ (primarily from natural gas systems with a negligible contribution from the oil sector) while the enteric fermentation sector also contributes about 30 Gg yr⁻¹. An additional 25 Gg yr⁻¹ comes from other sources. The implications of this are two-fold: (1) there is potentially an additional 55 Gg of CH₄ emissions annually impacting observations within the N/E wind sector at BAO that are not from the oil and gas industry, and thus, were not considered in the simple two-member mixing model used in the CFRPS or L12; and (2) the 71 Gg yr⁻¹ of CH₄ emissions estimated from bottom-up methods for the oil and gas industry in Weld county in the CFRPS are more than double the EDGAR inventory for this sector. This second point suggests that there is a problem with the base year EDGAR inventory and not with the prescribed 10% scaling term since the base year for both the CFRPS and the EDGAR inventory is 2008. Importantly, neither the EDGAR nor the CFRPS bottom-up inventories are wholly representative of CH₄ emissions in this region. In fact, as shown in Fig. 9, it is found that by combining the sectors not related to the oil and gas industry in the EDGAR inventory with the CFRPS inventory (which includes the oil and gas sector only), the total emissions in the bottom-up inventory are in relative agreement with the top-down

observations ($^{\sim}12\%$ overestimate in the bottom-up). It is worth noting that repeating this analysis for slightly larger or slightly smaller footprints gives consistent results, with bottom-up vs top-down estimates agreeing to within $\pm10\%$ when the CFRPS oil and gas emissions are merged with the EDGAR inventory.

In the Denver metro counties the natural gas systems sector contributes about 30-35% of total CH₄ emissions, which is similar to the contribution of natural gas systems to CH₄ emissions in Weld/Larimer counties. Scaling up the natural gas systems sector in the EDGAR inventory for the Denver metro counties, therefore, would result in a large over estimate of CH₄ emissions. A key difference, however, is the types of sources within the natural gas systems sector contributing to emissions in Denver vs Weld/Larimer. According to statistics about natural gas production in the region (COGCC, 2011), natural gas production in the Denver metro counties is about 4% of that in Weld county. Thus, emissions to the south and east of BAO likely arise from refineries and/or gas distribution networks from which the EDGAR bottom-up inventory estimates emissions that are consistent with the observations at BAO.

3.3.5 Benzene

Total benzene emissions in Weld/Larimer counties are estimated to be 0.64 (0.22-1.44) Gg yr $^{-1}$ and 0.95 (0.71 – 1.28) Gg yr $^{-1}$ in the Denver metro counties. Benzene emissions in the NEI08 inventory are 0.49 Gg yr $^{-1}$ for Weld/Larimer counties, at the low end of the observed uncertainty range; the Denver metro emissions are 1.30 Gg yr $^{-1}$, which is just outside the upper limit of the top-down uncertainty range. This suggests that the NEI08 benzene emissions are not inconsistent with the observations at 95% confidence, but would likely not be in agreement under relaxed confidence criteria. The top-down emissions from the CFRPS (0.39 - 1.19 Gg yr $^{-1}$) are consistent with our top-down estimates.

Mobile combustion and evaporative sources represent the largest contributor to benzene emissions in the NEI08 inventory in both the Denver metro counties and in Weld/Larimer counties. In the Denver metro counties mobile sector vehicles contribute to 92% of the benzene emissions, compared with 80% in Weld/Larimer combined and 50% in Weld County alone. Therefore, a comparison of top-down estimates to the bottom-up inventories of benzene from Weld County (and not over a larger footprint, as we have determined) may be biased. The CFRPS bottom-up estimate for benzene emissions, which corresponds to venting and flashing emissions in Weld county only, is 0.14 (0.05 - 0.23) Gg yr⁻¹. This equates to a significant fraction of what the NEI08 estimates for the region: 10 - 49% of the total Weld/Larimer benzene NEI08 emissions and 20-90% of the Weld County NEI08 emissions. According to the NEIO8 inventory, the non-combustion sources in the industrial sector, where the venting and flashing emissions would occur, account for only 11% of the total Weld/Larimer emissions and 20% of the Weld County emissions. It is likely, then, that the benzene sources of the CFRPS inventory and the NEIO8 inventory are mutually exclusive, and a better estimate for Weld/Larimer counties might be the sum of both inventories. The sum of the bottom-up CFRPS and NEI08 estimates for Weld/Larimer counties is 0.54 - 0.72 Gg yr⁻¹, in better agreement with the top-down estimates (both this study and the CFRPS) than either of the bottom-up inventories alone. Better agreement of the NEI08 benzene emissions with observations for the Denver metro region suggests that the mobile sources are wellaccounted for or even slightly over-estimated in the NEI08 inventory.

3.3.6 C_3 - C_5 alkanes

Absolute emissions of the 4 studied C_3 - C_5 alkanes in Weld/Larimer counties are comparable or greater than those estimated for the Denver metro counties. Propane and n-butane exhibit the largest enhancements, with total emissions in Weld County being 2.5 and 2.4 times higher, respectively, than that of the Denver metro counties. Emissions of *iso*-pentane, which is associated with gasoline vehicle emissions (Watson et al., 2001) in addition to oil and gas production (Pétron et al., 2012), are similar in the two regions, and total emissions of *n*-pentane, associated mainly with oil and gas production in the region (Watson et al., 2001; Pétron et al., 2012), are about 1.9 times higher in Weld/Larimer counties.

In the Denver metro counties the bottom-up estimates in the NEI 2005 inventory show a consistent over-estimate for most of the alkanes, relative to the top-down observations. The one exception is propane, which is significantly underestimated (by a factor of ~10) in the bottom-up inventory. We see better agreement in Weld/Larimer counties, but the same general relationship in the bottom-up vs top-down comparison: propane is underestimated in the NEI05 inventory relative to the observations (although it is within the lower uncertainty bracket), while the other alkanes are overestimated, but also within the uncertainty brackets. Recent emissions cutting regulations targeting the oil and gas industry in Colorado may explain the overestimate of the alkanes in the NEI05 bottom-up inventory; however, there appears to be a systematic underestimate of propane. There have been similar findings in two other recent studies that utilized $^{14}CO_2$ observations (Turnbull et al., 2011; Miller et al., 2012) to compare top-down and bottom-up estimates of propane.

A chemical sink of the C₃-C₅ alkanes would theoretically lead to an underestimate of emissions by the top-down methods used here, with stronger biases for the larger, shorter-lived alkanes. As detailed in Table 2, the lifetimes of these alkanes range from 3 days, for the pentanes, to 5 days for n-butane, to 12 days, for propane. There was some evidence in the CFRPS for OH chemistry influencing the ratios of various alkane tracers when measured close to sources vs at BAO (Pétron et al., 2012). In the east coast study of Miller et al. (2012), which used off-shore sampling sites well removed from local sources, summertime tracer/CO₂ff ratios were lower than wintertime ones, in proportion to their chemical lifetime with respect to OH. We find only minor differences (which are well within the uncertainties), however, in the relative ratios of alkanes with CO₂ff from summer to winter, so we use year-round observations in the current analysis. The seasonal impacts of photochemistry are likely minimized somewhat by subtracting a seasonally variable and relatively nearby background prior to calculating the enhancement ratios. For some perspective, an 8 hour atmospheric residence time at an average OH molecular density of 5 x 10⁶ cm⁻³ would result in an underestimate of about 10 Gg yr⁻¹ in the C₃H₈ emissions estimate, which is well within the uncertainty brackets of the top-down estimate of 14-64 Gg yr⁻¹. Thus, while we cannot rule out the influence of OH chemistry, we assume that any chemical effects are small relative to other sources of uncertainty in the analysis.

4. Implications for Carbon Monoxide and Methane Inventories

Observations of Δ^{14} CO₂ at the BAO tower have provided an important tracer with which to better constrain and evaluate emissions of a suite of compounds important to climate and air quality related to combustion and industrial processes along the northern Colorado Front Range. In this section, we will discuss important implications of our findings related to our evaluation of bottom-up inventories of carbon monoxide and methane, two globally important atmospheric trace gases.

4.1 Carbon Monoxide Bottom-up Inventory

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As discussed in Sec 3.3.2, our observations and evaluation of the NEI08 inventory are consistent with prior findings that CO emissions are overestimated at the national level in previous versions of the (Parrish, 2006; Hudman et al., 2008; Miller et al., 2008). There is evidence, albeit from a limited number of samples, that the California county-level bottom-up emissions of CO more accurately reflect the emissions estimated from atmospheric observations (Fig. 4). This provides the motivation to investigate whether there are fundamental differences in the methods for compiling the bottom-up CO inventory in California vs Colorado, as well as in other states.

An analysis of the mobile sector CO (NEI08) and CO₂ff (Vulcan08) emissions in comparison with the onroad observations of tail-pipe emissions of CO by Bishop and Stedman (2008) in Denver and Los Angeles suggests that NEI08 CO emissions in the mobile sector, specifically, are biased high in Colorado. Their observations show only very small differences between the CO-to-fuel-burnt ratio (and therefore the CO/CO₂ ratio) emitted from vehicles in Denver (in 2006) and Los Angeles (in 2008). The CO-to-fuel-burnt ratios observed in the two cities correspond to CO/CO₂ emission ratios of 16 ppb/ppm (Denver in 2006) and 18 ppb/ppm (Los Angeles in 2006). In Los Angeles, the observed ratio (Bishop and Stedman, 2008) closely resembles the bottom-up ratio of 17 ppb/ppm calculated from the NEI08 and Vulcan inventories for the mobile sector only; however, the mobile sector bottom-up ratio for Denver is 40 ppb/ppm, 2.5 times the observed ratio. Similarly, in Weld/Larimer counties the bottom-up ratio is 43 ppb/ppm and U.S.-wide it is 38 ppb/ppm. It should be noted that the mobile CO emissions in California are estimated using a different mobile source model, EMFAC2007 (EMFAC, hereafter) (CARB, 2007) than what is used for the rest of the United States, MOBILE6.2 (MOBILE, hereafter) (EPA, 2012a). In light of our comparison of CO/CO₂ff observations between California and other regions, and given the Bishop and Stedman (2008) observations in comparison with the mobile sector bottom up inventories, it appears likely that the MOBILE CO emission factor outputs are biased high relative to the EMFAC model.

This is also consistent with a recent comparison (Fujita et al., 2012) of these two mobile source models, along with the MOVES2010a (MOVES, hereafter) model, which was recently adopted by the EPA (EPA, 2010). This study showed that MOBILE emission factor outputs are biased high relative to the EMFAC and MOVES outputs, both of which showed close agreement with observations in a Los Angeles tunnel. This analysis showed that CO was over-estimated by the MOBILE model by a factor of 1.6-2.0 across a range of temperatures and traffic conditions and was found to be relatively insensitive to whether emission control programs were included in the model inputs. Both EMFAC and MOBILE use a region-wide average driving schedule and speed to compile emission factors for different vehicle types and model years, while MOVES uses a more specific approach, where emission factors are calculated for different speed and power bins. The consistency between MOVES, EMFAC, and the observations

detailed in Fujita et al. (2012), suggests that the general framework of the MOBILE model for scaling up emission factors from individual vehicles for an average driving schedule, which is essentially the same as that used by the EMFAC model, is not the issue. Rather, the emission ratios associated with individual vehicle types and model years needs to be adjusted. A separate study by the Federal Highway Administration Resource Center (Claggett and Houk, 2008) also analyzed differences in emission factor outputs by the EMFAC and MOBILE models and found that across a spectrum of average vehicle speeds for identical vehicle fleets in 2010 (a future scenario in the 2008 study), the MOBILE model CO emission factors were higher than the EMFAC output by 50%-300%, depending on the average vehicle speed. More recently, a study comparing CO and NO_x observations in Ada County, Idaho with outputs from both MOBILE and MOVES found differences in the CO/NO_x ratio outputs from the two source models, though most of the differences were attributed to differences in NO_x, not CO (Wallace et al., 2012).

With the EPA soon to adopt MOVES2010a for the NEI, and given the analysis of Fujita et al. (2012), future releases of the NEI inventory are predicted to produce more accurate estimates of CO emissions for Colorado and the rest of the U.S. However, continued evaluations of these mobile source models should be performed alongside observations such as those presented here, as well as those from Bishop and Stedman (2008) and Fujita et al. (2012).

We derive a modified bottom-up CO estimate for the Denver metro counties and Weld/Larimer counties in which the NEI08 mobile sector CO emission rate is replaced with a new estimate calculated from the Vulcan08 mobile sector CO₂ emission rate and the observed mobile sector CO/CO₂ ratio of 16 ppb/ppm from Bishop and Stedman (2008). This new estimate, shown in Fig. 6 and labeled as 'Modified NEI', brings the the top-down and bottom-up values to within 30% for the Denver metro counties and to within 10% for Weld/Larimer counties. The remaining discrepancy between these modified estimates and the observations could be a result of some combination of: (1) diesel vehicles that contribute significantly to the mobile sector CO₂ emissions but are not a significant source of CO; and (2) a reduction in the CO-to-fuel-burnt emission ratio between 2008 and 2009-2010. Additionally, this crude scaling exercise does not take into account changes in the CO/CO₂ emission ratio during "cold starts", which likely introduces significant variability at smaller spatial scales and shorter temporal scales. With these caveats acknowledged, scaling-up of this modified CO inventory reduces the total anthropogenic source of CO in the United States from 60 Tg yr⁻¹ to 39 Tg yr⁻¹, close to the 60% reduction recommended by Hudman et al. (2008) for anthropogenic CO emissions in the United States and close to the national ¹⁴CO₂ based estimate of 41 (33-53) Tg yr⁻¹ (Miller et al., 2012).

4.2 Methane from oil and gas production

The findings of the CFRPS suggested the presence of enhanced CH_4 levels at BAO over that predicted by bottom up statistical data on condensate tanks and raw natural gas profiles within the Denver Julesberg Basin (Pétron et al., 2012). These findings have since been scrutinized for not considering agricultural sources of CH_4 (Sgamma, 2012) and for the methodology of using methane-to-propane ratios to derive its top-down estimates (Levi, 2012). The results presented here bring a new perspective to this problem, and we arrive at conclusions that are largely independent of the assumptions and methodology used in the CFRPS and in L12. From our analysis of the EDGARv4.2 CH4 emissions estimates for the region, we

find that the top-down methodologies in both the L12 and CFRPS studies are flawed in that they fail to consider additional sources of CH₄ that are not directly related to the oil and gas industry in the region. We find that the bottom-up CH₄ emission estimate derived in the CFRPS is not inconsistent with the top-down estimates using observations at the BAO tower, but that additional sources, primarily enteric fermentation, must be also considered. In these previous studies the contribution from other sources to the CH₄ signal at BAO has resulted in either an overestimate of the venting/fugitive CH₄ source (CFRPS) or a larger methane-to-propane ratio for raw natural gas than is likely to be present in the region (L12).

Natural gas systems are a large source of CH₄ on continental and global scales according to the EDGAR inventory and others (e.g. (EPA, 2012b)). A comparison of the CFRPS bottom-up estimate and the oil and gas systems sector in the EDGARv4.2 inventory suggests that an additional ~40 Gg yr⁻¹ are missing from the EDGAR inventory for the region. This equates to an underestimate of about 60% for the oil and gas sector within the EDGAR inventory, on a local scale. One explanation for this underestimate could be an error in the spatial allocation of CH₄ emissions, such that emissions are estimated accurately over larger spatial scales but local inaccuracies arise when geographically distributing the emissions. Thus, it is impossible to speculate on whether this underestimate is systematic across the oil and gas sector in the EDGAR inventory; however, it should be noted that, in theory, this underestimate would scale up to a significant underestimate (~30%) of total CH₄ emissions across the United States. More observations are recommended, therefore, across a range of spatial and temporal scales in different regions to properly evaluate EDGAR, and other CH₄ emission databases, in the context of continental or global scale emissions, which could have potential implications for global atmospheric chemistry and climate. Our results provide strong motivation for the continued use of ¹⁴CO₂ observations in these evaluations.

5 Summary and Conclusions

We have analyzed 145 whole air samples for $\Delta^{14}\text{CO}_2$ collected across 15 months at the NOAA BAO tall tower in Erie, Colorado. Air sampled at this site is heavily impacted by emissions from a variety of sources including urban, rural, and industrial activities. The oil and gas industry, in particular, was found to contribute to enhancements in a number of industry-related trace gases relative to fossil fuel CO_2 when the tower is downwind of oil and gas activities, which are concentrated in Weld County to the north and east. The observed enhancements suggest that CH_4 , $\text{C}_3\text{-C}_5$ alkanes, and benzene emissions (relative to CO_2 ff emissions) are a factor of 5, ~10, and 1.6 greater, respectively, in air masses arriving passing over Weld and Larimer counties (north and east) over those originating from the Denver metro counties (south).

With the availability of a spatially resolved bottom-up CO_2 ff emissions data product from the Vulcan Project, we are able to take advantage of correlations of various trace gases with CO_2 ff, derived from $\Delta^{14}CO_2$ observations, in order to critically evaluate the accuracy of the bottom-up emissions inventories of these gases. The primary source of uncertainty in this approach is the uncertainty in the spatial extent of the observation footprint, and therefore the precise reference emissions value for CO_2 ff. This is particularly relevant for Weld and Larimer counties, which presents an extreme case where emissions related to the oil and gas industry are confined to a well-defined region within a larger region of significant CO_2 ff emissions from mobile sources and there is significant spatial heterogeneity in

tracer/CO₂ff emission ratios. Observations of trace gases alongside ¹⁴CO₂ observations on a mobile platform throughout the N/E sector (i.e. the sampling approach used in the CFRPS) would help to resolve this issue, as would the incorporation of an atmospheric transport model such as STILT (Lin et al., 2003) into the analysis. We find that our results for the Denver metro counties are less sensitive to assumptions regarding the spatial extent of the observation footprint, at least for CH₄. Within the constraints set by this and other sources of uncertainty, we conclude that our top-down estimates are inconsistent with the bottom up inventories for CH₄, CO, benzene, and propane.

CO is underestimated in the NEI08 inventory in both Weld/Larimer counties and the Denver metro counties by a factor of $^{\sim}2$, consistent with prior evaluations of earlier NEI inventories in the U.S. over larger scales. From the NEI08 CO and Vulcan08 CO₂ff emissions estimates, we calculate that the average emission factor from on-road gasoline vehicles is $^{\sim}40$ ppb CO/ppm CO₂ for the region, while our observations are more consistent with an emission ratio of 16 ppb/ppm for these vehicles.

For Weld/Larimer counties, we find that emissions of benzene and CH₄ are underestimated in the CFRPS bottom-up inventory, but that this underestimate can be explained by considering the contribution of emissions from other sectors (not related to the oil and gas industry) to the signal of these gases at the BAO. We conclude, therefore, that the CFRPS bottom-up CH₄ and benzene estimates are not inconsistent with top-down estimates from BAO, in contrast to the findings of Pétron et al. (2012). Our analysis of the CFRPS and other bottom-up emission estimates suggest that in Weld/Larimer counties the mobile sector contributes to about 75-80% of benzene emissions, enteric fermentation (e.g. cattle feedlots) contributes about 25% of CH₄ emissions, and other sources account for an additional 20% of CH₄ emissions. These results are consistent with 1.7% of total natural gas production in the DJB being vented to the atmosphere, lower than the range of 2.3-7.7% estimated by Pétron et al. (2012). Future studies of the impact of the oil and gas industry on atmospheric composition using observations at BAO need to consider these additional emission sources. Additional observations of ¹⁴CO₂ and CH₄ in other oil and gas producing regions and comparisons to bottom-up inventories are recommended to determine whether these results are specific to the DJB or if they are more general.

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References

Andrews, A. E., Kofler, J. D., Trudeau, M. E., Williams, J. C., Neff, D. H., Masaire, K. A., Chao, D. Y., Kitzis, D. R., Novelli, P. C., Zhao, C. L., Dlugokencky, E. J., Lang, P. M., Crotwell, M. J., Fischer, M. L., Parker, M. J.,

- Lee, J. T., Baumann, D. D., Desai, A. R., Stanier, C. O., de Wekker, S. F. J., Wolfe, D. E., Munger, J. W., and Tans, P. P.: Atmos. Meas. Technol. Discuss., submitted.
- Atkinson, R., Baulch, D. L., Cox, R. A., Crowley, J. N., Hampson, R. F., Hynes, R. G., Jenkin, M. E., Rossi, M.
- J., and Troe, J.: Evaluated kinetic and photochemical data for atmospheric chemistry: Volume II gas
- phase reactions of organic species, Atmos. Chem. Phys., 6, 3625-4055, 2006.

1088

1091

1095

1098

1101

1103

1107

1110

1113

1116

1119

1122

1125

- Bishop, G. A., and Stedman, D. H.: A decade of on-road emissions measurements, Environ. Sci. Technol., 42, 1651-1656, 2008.
- 1092 Brennan, P. J.: Los Angeles Wildfire Sends Pollution as Far as Denver, Bloomberg,
- http://www.bloomberg.com/apps/news?pid=newsarchive&sid=aJrSkTFs1dL4, access: March 7, 2012, 2009.
- 1096 California Air Resources Board Mobile Sources Emissions Inventory:
- 1097 (http://www.arb.ca.gov/msei/onroad/latest_version.htm) access: October 10, 2007.
- 1099 Claggett, M., and Houk, J.: Comparing MOBILES6.2 and Emfac2007 Emission Factors, Transp. Res. 1100 Record, 51-57, 2008.
- 1102 Colorado Oil and Gas Information System: Data Base accessible at http://cogcc.state.co.us, 2011.
- 104 Conway, T. J., Tans, P. P., Waterman, L. S., and Thoning, K. W.: Evidence for Interannual Variability of the 1105 Carbon-Cycle from the National-Oceanic-and-Atmospheric-Administration Climate-Monitoring-and-1106 Diagnostics-Laboratory Global-Air-Sampling-Network, J. Geophys. Res. Atmos., 99, 22831-22855, 1994.
- 1108 Currie, K. I., Brailsford, G., Nichol, S., Gomez, A., Sparks, R., Lassey, K. R., and Riedel, K.: Tropospheric (CO2)-C-14 at Wellington, New Zealand: the world's longest record, Biogeochem., 104, 5-22, 2011.
- Djuricin, S., Pataki, D. E., and Xu, X. M.: A comparison of tracer methods for quantifying CO(2) sources in an urban region, J. Geophys. Res.-Atmos., 115, 2010.
- Dlugokencky, E. J., Steele, L. P., Lang, P. M., and Masarie, K. A.: The Growth-Rate and Distribution of Atmospheric Methane, J. Geophys. Res. Atmos., 99, 17021-17043, 1994.
- 1117 U.S. Energy Information Administration CO2 Emissions Data:
- http://www.eia.gov/environment/data.cfm#summary, access: June 18, 2012, 2012.
- EPA: Official release of the MOVES2010 motor vehicle emissions model for emission inventories in SIPs and tranportation conformity., Fed. Regist., 75, 9411-9414, 2010.
- 1123 EPA Speciate Software Download Site: http://www.epa.gov/ttnchie1/software/speciate/, access: June 19, 2011.
- MOBILE6 Vehicle Emission Modeling Software: http://www.epa.gov/oms/m6.htm, access: October 10, 2012a.

- 1129 EPA: Inventory of U.S. Greenhouse Gas Emissions and Sinks: 1990-2010,
- 1130 http://www.epa.gov/climatechange/Downloads/ghgemissions/US-GHG-Inventory-2012-Main-Text.pdf,
- 1131 access: September 11, 2012, 2012b.
- 1132
- Fujita, E. M., Campbell, D. E., Zielinska, B., Chow, J. C., Lindhjem, C. E., DenBleyker, G. A., Bishop, G. A.,
- 1134 Schuchmann, B. G., Stedman, D. H., and Lawson, D. R.: Comparison of the MOVES2010a, MOBILE6.2, and
- 1135 EMFAC2007 mobile source emission models with on-road traffic tunnel and remote sensing
- measurements, J. Air Waste Mgmt. Assoc., 62, 1134-1149, doi:10.1080/10962247.2012.699016, 2012.
- 1137
- 1138 Godwin, H.: Half-Life of Radiocarbon, Nature, 195, 984-&, 1962.
- 1139
- 1140 Graven, H. D., Guilderson, T. P., and Keeling, R. F.: Methods for high-precision C-14 AMS measurement
- of atmospheric CO2 at LLNL, Radiocarbon, 49, 349-356, 2007.
- 1142
- Graven, H. D., Stephens, B. B., Guilderson, T. P., Campos, T. L., Schimel, D. S., Campbell, J. E., and Keeling,
- 1144 R. F.: Vertical profiles of biospheric and fossil fuel-derived CO(2) and fossil fuel CO(2): CO ratios from
- airborne measurements of Delta(14)C, CO(2) and CO above Colorado, USA, Tellus B, 61, 536-546, 2009.
- 1146
- 1147 Graven, H. D., and Gruber, N.: Continental-scale enrichment of atmospheric 14CO2 from the nuclear
- power industry: potential impact on the estimation of fossil fuel-derived CO2, Atmos. Chem. Phys., 11,
- 1149 12339-12349, 10.5194/acp-11-12339-2011, 2011.
- 1150
- Graven, H. D., Guilderson, T. P., and Keeling, R. F.: Observations of radiocarbon in CO2 at seven global
- sampling sites in the Scripps flask network: Analysis of spatial gradients and seasonal cycles J. Geophys.
- 1153 Res., 117, doi:10.1029/2011JD016535, 2012a.
- 1154
- 1155 Graven, H. D., Guilderson, T. P., and Keeling, R. F.: Observations of radiocarbon in CO2 at La Jolla,
- 1156 California, USA 1992-2007: Analysis of the long-term trend, J. Geophys. Res., 117,
- 1157 doi:10.1029/2011JD016533, 2012b.
- 1158
- Griffin, R. J., Chen, J. J., Carmody, K., Vutukuru, S., and Dabdub, D.: Contribution of gas phase oxidation
- of volatile organic compounds to atmospheric carbon monoxide levels in two areas of the United States,
- 1161 J. Geophys. Res.-Atmos., 112, doi: 10.1029/2006jd007602, 2007.
- 1162
- 1163 Gurney, K. R., Mendoza, D. L., Zhou, Y. Y., Fischer, M. L., Miller, C. C., Geethakumar, S., and Du Can, S. D.:
- High Resolution Fossil Fuel Combustion CO(2) Emission Fluxes for the United States, Environ. Sci.
- 1165 Technol., 43, 5535-5541, 2009.
- 1166
- Gurney, K. R., Chandrasekaran, V., Mendoza, D. L., and Geethakumar, S.: Quantification of uncertainty
- associated with NACP high resolution fossil fuel CO2 emissions: updates, challenges and future plans,
- 1169 North American Carbon Program All-Inverstigators Meeting, New Orleans, LA, 2011.
- 1170
- Hsueh, D. Y., Krakauer, N. Y., Randerson, J. T., Xu, X. M., Trumbore, S. E., and Southon, J. R.: Regional
- patterns of radiocarbon and fossil fuel-derived CO2 in surface air across North America, Geophys. Res.
- 1173 Lett., 34, 2007.
- 1174
- Hudman, R. C., Jacob, D. J., Cooper, O. R., Evans, M. J., Heald, C. L., Park, R. J., Fehsenfeld, F., Flocke, F.,
- Holloway, J., Hubler, G., Kita, K., Koike, M., Kondo, Y., Neuman, A., Nowak, J., Oltmans, S., Parrish, D.,

- 1177 Roberts, J. M., and Ryerson, T.: Ozone production in transpacific Asian pollution plumes and implications
- 1178 for ozone air quality in California, J. Geophys. Res., 109, doi:10.1029/2004JD004974, 2004.
- 1179
- Hudman, R. C., Murray, L. T., Jacob, D. J., Millet, D. B., Turquety, S., Wu, S., Blake, D. R., Goldstein, A. H.,
- Holloway, J., and Sachse, G. W.: Biogenic versus anthropogenic sources of CO in the United States,
- 1182 Geophys. Res. Lett., 35, 2008.
- 1183
- Lee, B. H., Munger, J. W., Wofsy, S. C., and Goldstein, A. H.: Anthropogenic emissions of nonmethane
- hydrocarbons in the northeastern United States: Measured seasonal variations from 1992-1996 and
- 1186 1999-2001, J. Geophys. Res.-Atmos., 111, 2006.
- 1187
- 1188 Lehman, S. J., Miller, J. B., Wolak, C., Southon, J., Tans, P. P., Montzka, S. A., Sweeney, C., Andrews, A.,
- LaFranchi, B. W., Guilderson, T. P., Fischer, M. L., and Turnbull, J. C.: Allocation of terrestrial carbon
- sources using 14CO2; measurement and modeling, Radiocarbon, submitted.
- 1191
- Levi, M. A.: Comment on "Hydrocarbon emissions characterization in the Colorado Front Range A Pilot
- 1193 Study", J. Geophys. Res. Atmos., in press, 2012.
- 1194
- 1195 Levin, I., Kromer, B., Schmidt, M., and Sartorius, H.: A novel approach for independent budgeting of
- fossil fuel CO2 over Europe by (CO2)-C-14 observations, Geophys. Res. Lett., 30, 2003.
- 1197
- Levin, I., and Kromer, B.: The tropospheric (CO2)-C-14 level in mid-latitudes of the Northern Hemisphere
- 1199 (1959-2003), Radiocarbon, 46, 1261-1272, 2004.
- 1200
- Levin, I., and Karstens, U. T. E.: Inferring high-resolution fossil fuel CO2 records at continental sites from
- combined 14CO2 and CO observations, Tellus B, 59, 245-250, doi: 10.1111/j.1600-0889.2006.00244.x,
- 1203 2007.
- 1204
- 1205 Levin, I., Naegler, T., Kromer, B., Diehl, M., Francey, R. J., Gomez-Pelaez, A. J., Steele, L. P., Wagenbach,
- 1206 D., Weller, R., and Worthy, D. E.: Observations and modelling of the global distribution and long-term
- trend of atmospheric 14CO(2), Tellus B, 62, 26-46, 2010.
- 1208
- Lin, J. C., Gerbig, C., Wofsy, S. C., Andrews, A. E., Daube, B. C., Davis, K. J., and Grainger, C. A.: A near-
- 1210 field tool for simulating the upstream influence of atmospheric observations: The Stochastic Time-
- 1211 Inverted Lagrangian Transport (STILT) model, J. Geophys. Res. Atmos., 108, 2003.
- 1212
- Miller, J. B., Lehman, S. J., Montzka, S. A., Sweeney, C., Miller, B. R., Wolak, C., Dlugokencky, E. J.,
- 1214 Southon, J., Turnbull, J. C., and Tans, P. P.: Linking emissions of fossil fuel CO2 and other anthropogenic
- trace gases using atmospheric 14CO2, J. Geophys. Res. Atmos., doi:10.1029/2011JD017048, 2012.
- 1216
- Miller, S. M., Matross, D. M., Andrews, A. E., Millet, D. B., Longo, M., Gottlieb, E. W., Hirsch, A. I., Gerbig,
- 1218 C., Lin, J. C., Daube, B. C., Hudman, R. C., Dias, P. L. S., Chow, V. Y., and Wofsy, S. C.: Sources of carbon
- 1219 monoxide and formaldehyde in North America determined from high-resolution atmospheric data,
- 1220 Atmos. Chem. Phys., 8, 7673-7696, 2008.
- 1221
- 1222 Montzka, S. A., Myers, R. C., Butler, J. H., Elkins, J. W., and Cummings, S. O.: Global Tropospheric
- 1223 Distribution and Calibration Scale of Hcfc-22, Geophys. Res. Lett., 20, 703-706, 1993.
- 1224

- 1225 Muller, S. A., Joos, F., Plattner, G. K., Edwards, N. R., and Stocker, T. F.: Modeled natural and excess
- 1226 radiocarbon: Sensitivities to the gas exchange formulation and ocean transport strength, Global
- 1227 Biogeochem. Cy., 22, 2008.

- 1229 Chemical Kinetics and Photochemical Data for Use in Atmospheric Studies Evaluation Number 15:
- 1230 http://jpldataeval.jpl.nasa.gov/, 2006.

1231

1232 National Emissions Inventory: http://www.epa.gov/ttn/chief/eiinformation.html, 2008.

1233

- 1234 Novelli, P. C., Connors, V. S., Reichle, H. G., Anderson, B. E., Brenninkmeijer, C. A. M., Brunke, E. G.,
- 1235 Doddridge, B. G., Kirchhoff, V. W. J. H., Lam, K. S., Masarie, K. A., Matsuo, T., Parrish, D. D., Scheel, H. E.,
- and Steele, L. P.: An internally consistent set of globally distributed atmospheric carbon monoxide
- mixing ratios developed using results from an intercomparison of measurements, J. Geophys. Res.
- 1238 Atmos., 103, 19285-19293, 1998.

1239

- Parrish, D. D.: Critical evaluation of US on-road vehicle emission inventories, Atmos. Environ., 40, 2288-
- 1241 2300, 2006.

1242

- 1243 Pétron, G., Frost, G., Hirsch, A. I., Montzka, S. A., Karion, A., Miller, B. R., Trainer, M., Sweeney, C.,
- 1244 Andrews, A. E., Miller, L., Kofler, J., Ryerson, T., Patrick, L., Siso, C., Kolodzey, W., Lang, P., Dlugokencky,
- 1245 E., Conway, T., Novelli, P., Masarie, K., Hall, B., Guenther, D., Kitzis, D., Miller, J. B., Neff, W., Wolfe, D.,
- 1246 and Tans, P. P.: Hydrocarbon emissions characterization in the Colorado front range A pilot study,
- 1247 117, doi:10.1029/2011JD016360, 2012.

1248

- Schuur, E. A. G., Trumbore, S. E., Mack, M. C., and Harden, J. W.: Isotopic composition of carbon dioxide
- 1250 from a boreal forest fire: Inferring carbon loss from measurements and modeling, Global Biogeochem.
- 1251 Cy., 17, 2003.

1252

- 1253 Sgamma, K. M.: Fracking: Colorado methane study not clear-cut, Nature, 483, 407-407,
- 1254 doi:10.1038/483407e, 2012.

1255

- 1256 Colorado Active Gas and Oil Wells April 2008: http://skytruth.mediatools.org/node/9590, access:
- 1257 October 1, 2008.

1258

1259 Stuiver, M., and Polach, H. A.: Reporting of C-14 Data - Discussion, Radiocarbon, 19, 355-363, 1977.

1260

1261 Suess, H. E.: Radiocarbon Concentration in Modern Wood, Science, 122, 415-417, 1955.

1262

- Sweeney, C., Gloor, E., Jacobson, A. R., Key, R. M., McKinley, G., Sarmiento, J. L., and Wanninkhof, R.:
- 1264 Constraining global air-sea gas exchange for CO2 with recent bomb (14)C measurements, Global
- 1265 Biogeochem. Cy., 21, 2007.

1266

- 1267 Thompson, M. V., and Randerson, J. T.: Impulse response functions of terrestrial carbon cycle models:
- method and application, Glob. Change Biol., 5, 371-394, 1999.

1269

- 1270 Thoning, K. W., Tans, P. P., and Komhyr, W. D.: Atmospheric Carbon-Dioxide at Mauna Loa Observatory
- 1271 .2. Analysis of the NOAA GMCC Data, 1974-1985, J. Geophys. Res. Atmos., 94, 8549-8565, 1989.

- 1273 Turnbull, J., Rayner, P., Miller, J., Naegler, T., Ciais, P., and Cozic, A.: On the use of (14)CO(2) as a tracer
- for fossil fuel CO(2): Quantifying uncertainties using an atmospheric transport model, J. Geophys. Res.
- 1275 Atmos., 114, 2009.

- Turnbull, J. C., Miller, J. B., Lehman, S. J., Tans, P. P., Sparks, R. J., and Southon, J.: Comparison of (CO2)-
- 1278 C-14, CO, and SF6 as tracers for recently added fossil fuel CO2 in the atmosphere and implications for
- biological CO2 exchange, Geophys. Res. Lett., 33, 2006.

1280

- Turnbull, J. C., Lehman, S. J., Miller, J. B., Sparks, R. J., Southon, J. R., and Tans, P. P.: A new high
- 1282 precision (14)CO(2) time series for North American continental air, J. Geophys. Res. Atmos., 112, 2007.

1283

Turnbull, J. C., Lehman, S. J., Morgan, S., and Wolak, C.: A New Automated Extraction System for (14)C Measurement for Atmospheric Co(2), Radiocarbon, 52, 1261-1269, 2010.

1285 1286

- Turnbull, J. C., Karion, A., Fischer, M. L., Faloona, I., Guilderson, T., Lehman, S. J., Miller, B. R., Miller, J.
- B., Montzka, S., Sherwood, T., Saripalli, S., Sweeney, C., and Tans, P. P.: Assessment of fossil fuel carbon
- 1289 dioxide and other anthropogenic trace gas emissions from airborne measurements over Sacramento,
- 1290 California in spring 2009, Atmos. Chem. Phys., 11, 705-721, 2011.

1291

- 1292 Van der Laan, S., Karstens, U., Neubert, R. E. M., Van der Laan-Luijkx, I. T., and Meijer, H. A. J.:
- 1293 Observation-based estimates of fossil fuel-derived CO2 emissions in the Netherlands using Delta 14C, CO
- 1294 and 222Radon, Tellus B, 62, 389-402, 2010.

1295

- Vaughn, B. H., Miller, J. B., Ferretti, D. F., and White, J. C.: Stable isotope measurements of atmospheric
- 1297 CO2 and CH4, in: Handbook of Stable Isotope Analytical Techniques, edited by: de Groot, P., Elsevier
- 1298 Inc., San Diego, CA, 272-304, 2004.

1299

- 1300 Vay, S. A., Choi, Y., Vadrevu, K. P., Blake, D. R., Tyler, S. C., Wisthaler, A., Hecobian, A., Kondo, Y., Diskin,
- 1301 G. S., Sachse, G. W., Woo, J. H., Weinheimer, A. J., Burkhart, J. F., Stohl, A., and Wennberg, P. O.:
- 1302 Patterns of CO(2) and radiocarbon across high northern latitudes during International Polar Year 2008, J.
- 1303 Geophys. Res. Atmos., 116, 2011.

1304

- 1305 Wallace, H. W., Jobson, B. T., Erickson, M. H., McCoskey, J. K., VanReken, T. M., Lamb, B. K., Vaughan, J.
- 1306 K., Hardy, R. J., Cole, J. L., Strachan, S. M., and Zhang, W.: Comparison of wintertime CO to NOx ratios to
- 1307 MOVES and MOBILE6.2 on-road emissions inventories, Atmos. Environ., 63, 289-297,
- 1308 doi:10.1016/j.atmosenv.2012.08.062, 2012.

1309

- 1310 Warneke, C., McKeen, S. A., de Gouw, J. A., Goldan, P. D., Kuster, W. C., Holloway, J. S., Williams, E. J.,
- Lerner, B. M., Parrish, D. D., Trainer, M., Fehsenfeld, F. C., Kato, S., Atlas, E. L., Baker, A., and Blake, D. R.:
- 1312 Determination of urban volatile organic compound emission ratios and comparison with an emissions
- database, J. Geophys. Res.-Atmos., 112, 2007.

1314

- Watson, J. G., Chow, J. C., and Fujita, E. M.: Review of volatile organic compound source apportionment
- 1316 by chemical mass balance, Atmos. Environ., 35, 1567-1584, 2001.

1317

- 1318 Zondervan, A., and Meijer, H. A. J.: Isotopic characterisation of CO2 sources during regional pollution
- events using isotopic and radiocarbon analysis, Tellus B, 48, 601-612, 1996.

| | Wind | | | Taranto esta de | | Ratio co | onfidence limits | Bottom-Up | | | | | | Scaled | | Emissions | | | | |
|---------|----------|------|----|-----------------|------------|----------|------------------|-----------|-----------|-----------|------|-------|-------|-----------|-------|-----------|------------|----------------------------|-------|-------|
| Species | Sector | r2 | n | Rat | io (units) | | (2σ) | Emissions | Source | Base Year | α | | n/max | Emissions | | /Max | Top-Dowm E | missions (E _x) | | n/Max |
| | | | | | | min | max | | | | | min | max | | min | max | | | min | max |
| CO | N/E | 0.75 | 44 | 9.3 | (ppb/ppm) | 8.1 | 10.6 | 116.0 Gg | NEI08 | 2008 | 3.6 | -10.5 | 11 | 120.1 Gg | 103.8 | 128.4 | 65.8 | Gg | 50.1 | 81.7 |
| | S | 0.85 | 25 | 10.0 | (ppb/ppm) | 6.7 | 13.3 | 362.1 Gg | NEI08 | 2008 | 1.7 | -10.5 | 5 | 368.2 Gg | 324.1 | 380.5 | 175.1 | Gg | 106.8 | 241.2 |
| | Combined | 0.81 | 68 | 9.4 | (ppb/ppm) | 8.3 | 10.8 | 478.1 Gg | NEI08 | 2008 | | | | 488.3 Gg | 427.9 | 508.9 | 229.1 | Gg | 176.8 | 286.9 |
| CH4 | N/E | 0.87 | 44 | 30.0 | (ppb/ppm) | 24.1 | 33.3 | 64.3 Gg | CFRPS | 2009 | 10 | 0 | 30 | 70.7 Gg | 46.0 | 99.8 | 120.7 | Gg | 40.4 | 272.2 |
| | S | 0.69 | 26 | 9.8 | (ppb/ppm) | 6.0 | 13.3 | 97.7Gg | Edgar4.2 | 2008 | 1.7 | 0 | 5 | 99.3 Gg | 97.7 | 102.6 | 98.1 | Gg | 55.2 | 137.6 |
| C2H2 | N/E | 0.80 | 39 | 43.3 | (ppt/ppm) | 36.2 | 59.3 | 0.172 Gg | NEI05 | 2005 | 11.6 | 0 | 35 | 0.192 Gg | 0.172 | 0.232 | 0.284 | Gg | 0.210 | 0.403 |
| | S | 0.83 | 24 | 47.9 | (ppt/ppm) | 37.0 | 58.0 | 0.544 Gg | NEI05 | 2005 | 5.2 | 0 | 16 | 0.572 Gg | 0.544 | 0.629 | 0.776 | Gg | 0.541 | 1.001 |
| | Combined | 0.81 | 63 | 43.7 | (ppt/ppm) | 38.3 | 55.4 | 0.687 Gg | NEI05 | 2005 | - | | | 0.764 Gg | 0.716 | 0.861 | 0.995 | Gg | 0.760 | 1.326 |
| BENZ | N/E | 0.85 | 39 | 32.3 | (ppt/ppm) | 27.1 | 38.9 | 0.468 Gg | NEI08 | 2008 | 3.6 | 0 | 11 | 0.485 Gg | 0.468 | 0.518 | 0.64 | Gg | 0.22 | 1.44 |
| | S | 0.78 | 24 | 19.5 | (ppt/ppm) | 16.6 | 25.0 | 1.28 Gg | NEI08 | 2008 | 1.7 | 0 | 5 | 1.30 Gg | 1.28 | 1.34 | 0.95 | Gg | 0.71 | 1.28 |
| iC5H12 | N/E | 0.78 | 39 | 317.2 | (ppt/ppm) | 243.3 | 456.0 | 4.4 Gg | CFRPS | 2009 | 10 | 0 | 30 | 4.8 Gg | 1.0 | 9.6 | 5.8 | Gg | 1.9 | 13.4 |
| | S | 0.83 | 23 | 92.7 | (ppt/ppm) | 63.1 | 116.1 | 5.96 Gg | NEI05 | 2005 | 5.2 | 0 | 15.6 | 6.3 Gg | 6.0 | 6.9 | 4.2 | Gg | 2.6 | 5.5 |
| nC5H12 | N/E | 0.78 | 39 | 332.5 | (ppt/ppm) | 238.2 | 487.0 | 3.4 Gg | CFRPS | 2009 | 10 | 0 | 30 | 3.7 Gg | 1.0 | 8.5 | 6.0 | Gg | 1.8 | 14.1 |
| | S | 0.76 | 23 | 70.2 | (ppt/ppm) | 47.7 | 98.8 | 4.52 Gg | NEI05 | 2005 | 5.2 | 0 | 15.6 | 4.8 Gg | 4.5 | 5.2 | 3.1 | Gg | 2.0 | 4.6 |
| nC4H10 | N/E | 0.75 | 39 | 1126.4 | (ppt/ppm) | 764.4 | 1497.7 | 11.4 Gg | CFRPS | 2009 | 10 | 0 | 30 | 12.5 Gg | 5.0 | 23.7 | 16.5 | Gg | 4.7 | 37.7 |
| | S | 0.70 | 23 | 191.6 | (ppt/ppm) | 104.3 | 237.9 | 9.7 Gg | NEI05 | 2005 | 5.2 | 0 | 15.6 | 10.2 Gg | 9.7 | 11.2 | 6.9 | Gg | 3.5 | 9.1 |
| C3H8 | N/E | 0.76 | 39 | 2529.7 | (ppt/ppm) | 1758.6 | 3462.3 | 26.1 Gg | CFRPS | 2009 | 10 | 0 | 30 | 28.7 Gg | 15.0 | 55.8 | 28.0 | Gg | 8.3 | 64.6 |
| | S | 0.55 | 23 | | (ppt/ppm) | 220.9 | 496.8 | 1.2 Gg | NEI05 | 2005 | 5.2 | 0 | 15.6 | 1.3 Gg | 1.2 | 1.4 | 11.3 | Gg | 5.6 | 14.5 |
| CO2 | N/E | 220 | | | 1 | 22 | - | 2.94 Tg C | Vulcan2.0 | 2002 | 2.8 | | - | 3.02 Tg C | 2.42 | 3.63 | 3.60 | Tg C | 1.10 | 6.80 |
| | S | | | 1 | 1 | | | 7.27 Tg C | Vulcan2.0 | 2002 | 2.8 | | | 7.47 Tg C | 5.98 | 8.97 | | | | |

Table 1 Summary of top-down and bottom-up annual emissions for each tracer discussed in our analysis. Included in this table are: correlation coefficients for each tracer with CO₂ff, observed tracer/CO₂ff ratios, bottom-up emissions directly from each inventory and also scaled to the observation period, and estimated top-down emissions. Bottom-up emissions for CO₂ff are also summarized. Uncertainties on the scaled bottom-up emissions and the top-down emissions are described in Sec. 3.3 and Sec. 3.3.1. Top-down emissions for CO₂ff for the N/E wind sector are estimated using the top-down CH₄ estimates from the CFRPS and L12 as a quantitative reference, as described in Sec. 3.3.1. Bottom-up CH₄ estimates for the S wind sector are detailed in Fig. 8 and associated text.

| Species | Atmospheric Lifetime ^a | Major sources near BAO | | | | | |
|---|-----------------------------------|--|--|--|--|--|--|
| carbon monoxide (CO) | 49 days | on-road and non-road gasoline combustion | | | | | |
| methane (CH ₄) | 6.9 years | oil and gas systems ^b , waste water treatment, landfills, cattle feed lot | | | | | |
| acetylene (C ₂ H ₂) | 17 days | mobile sources (combustion) | | | | | |
| benzene (C ₆ H ₆) | 10 days | mobile sources (combustion and evaporative), oil and gas systems | | | | | |
| iso-pentane (i-C ₅ H ₁₂) | 3.0 days | mobile sources (combustion and evaporative), oil and gas systems | | | | | |
| n-pentane (n -C ₅ H ₁₂) | 3.1 days | oil and gas systems | | | | | |
| n-butane (n-C ₄ H ₁₀) | 4.9 days | oil and gas systems | | | | | |
| propane (C₃H ₈) | 12 days | oil and gas systems | | | | | |

Table 2 Summary of trace gas lifetimes and major emission sources influencing observations at BAO (Watson et al., 2001; Pétron et al., 2012). ^a Atmospheric lifetimes estimated for [OH] = 1 x 10⁶ cm⁻³ using published rate constant data (Atkinson et al., 2006; NASA, 2006). ^b Sources include condensate tanks, well drilling and completion, distribution systems, refineries.

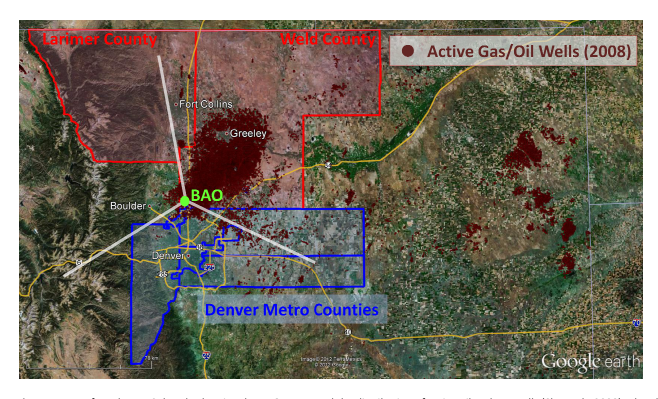


Figure 1 Map of northwest Colorado showing the BAO tower and the distribution of active oil and gas wells (SkyTruth, 2008). Also shown are the three wind sectors used to filter the data set for emission estimates in Weld/Larimer counties (North and East) and in the Denver metro counties (South). The Denver metro counties include Denver, Broomfield, Adams, Arapahoe, and Jefferson.

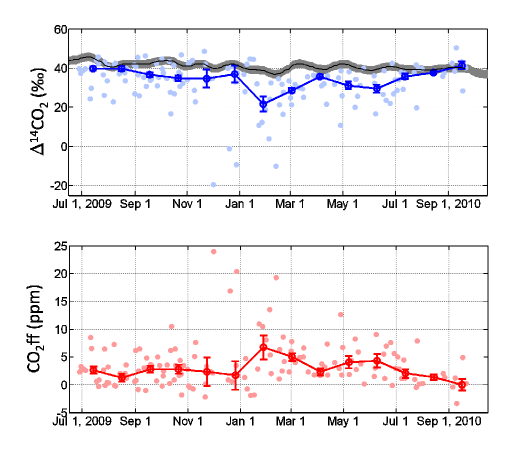


Figure 2 Time series of $^{14}CO_2$ (a) and CO_2 ff (b) from 145 discrete whole air samples (filled circles) collected at the BAO tower. Uncertainty in each $^{14}CO_2$ measurement is $\pm 2.2\%$, which translates to an uncertainty in each CO_2 ff observation of 1.2 ppm (see Sec. 3.1). Thirty day binned medians are shown as open circles in both (a) and (b), with error bars representing the standard error of the mean (1 σ) for each 30 day bin. Also shown in (a) is the $^{14}CO_2$ background as observed at NWR (black line) (Turnbull et al., 2007; Lehman et al., submitted), with the uncertainty envelope represented by the grey shaded region.

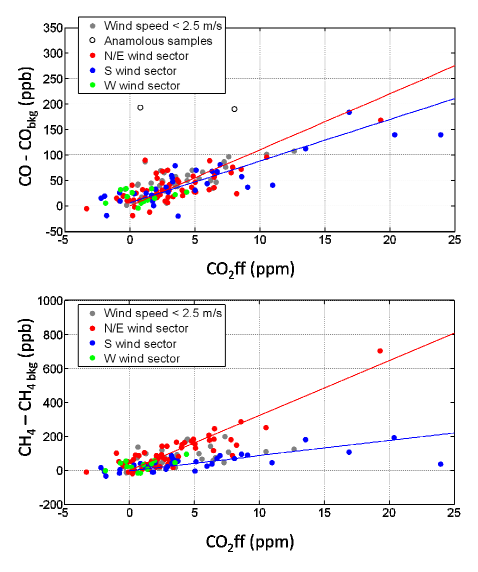


Figure 3 Correlation plots of CO (a) and CH4 (b) enhancements (with respect to background observations) with CO2ff. Data are separated into one of three wind sectors (north and east: red; south: blue; and west green), except in cases where average wind speeds were below 2.5 m/s over the 30 minutes prior to sampling. Best-fit lines are shown for the N/E and S wind sectors (correlation coefficients are given in Table 1). In (a), two points are shown as open circles which are omitted from our analysis (see Sec. 2.5).

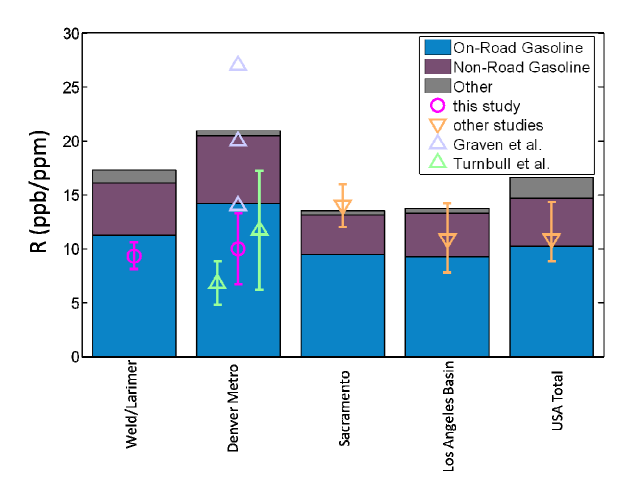


Figure 4 A comparison of CO/CO₂ff ratios observed or estimated in various US locations. The bars are calculated from bottom-up emissions estimates (NEI08 CO and Vulcan2.2 CO₂) and color-coded by the contribution of different sectors to the total CO emissions: on-road gasoline, non-road gasoline, and other. Observations from each location are shown, including those from our observations at BAO (split into Weld/Larimer and Denver metro influence based on wind sector) and observations from other studies: Denver (Turnbull et al., 2006; Graven et al., 2009)), Sacramento (Turnbull et al., 2011), LA Basin (which includes Los Angeles, Riverside, Orange, and San Bernardino counties) (Djuricin et al., 2010), and for the continental US (Miller et al., 2012).

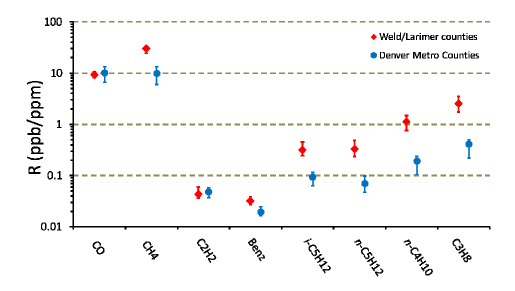


Figure 5 Observed tracer/CO₂ff ratios from Weld County (N/E wind sector, red diamonds) and the Denver metro counties (S wind sector, blue circles). Ratios are calculated as the median of the point-by-point ratios for all data where CO₂ff was detected above 1.2 ppm, as described in Sec. 3.3. Uncertainties in the median ratios are the 95% confidence intervals, defined as the 2.5-97.5 percentile range (~2σ confidence) from a distribution of 500 median estimates from a randomized re-sampling of the data (boot-strapping with replacement). Note that the figure is presented using a logarithmic scale.

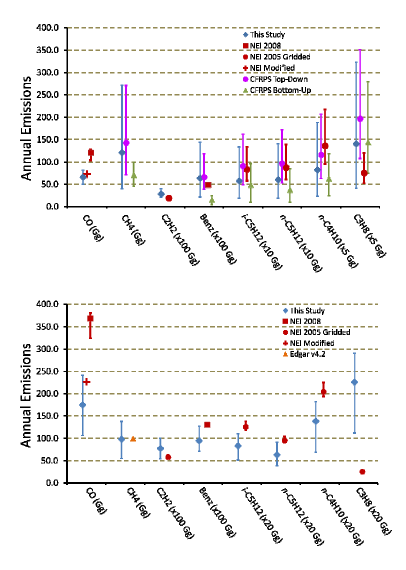


Figure 6 Emissions estimates from Weld County (a) and the Denver metro counties (b). Top-down emissions, calculated using Eq. 4, are shown as blue diamonds, with uncertainties given as described in Sec. 3.3. Bottom-up emissions estimates are included for comparison for each species, making use of NEI, EDGAR, and CFRPS (Pétron et al., 2012) inventories. Also shown are the top-down estimates from the CFRPS for CH₄, Benzene, and the alkanes in Weld County. The CFRPS error bars (both bottom-up and top-down) are related to uncertainties in the VOC profiles for specific emission sources related to the oil and gas industry in Weld County. Note the differences in units for the different trace gases.

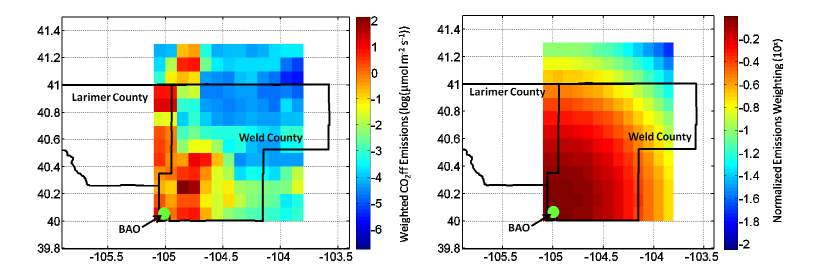


Figure 7 Weighted Vulcan CO₂ff emissions (scaled to 2009-2010) (a) and weighting function (b) for a theoretical observation footprint for N/E wind sector observations.

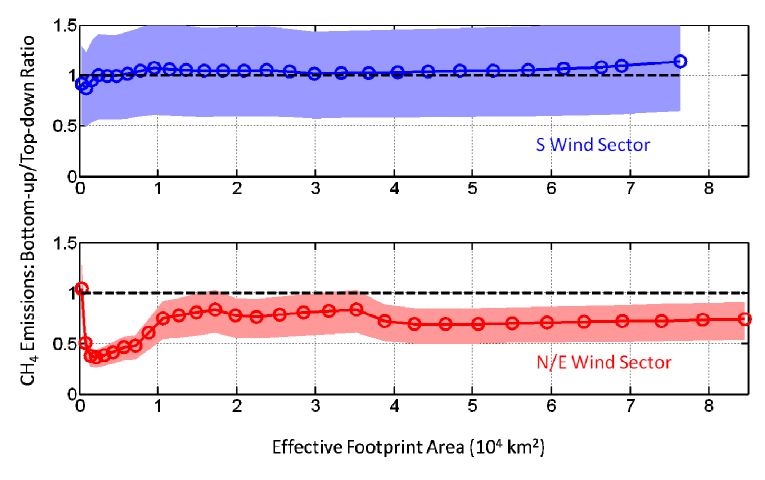


Figure 8 Ratio of bottom-up (EDGARv4.2) to top-down emissions estimates across an expanding hypothetical footprint for the South wind sector (top) and the North/East wind sector (bottom). A value of one means perfect agreement between bottom-up and top-down estimates. Shaded areas reflect the uncertainties in top-down emissions estimates only.

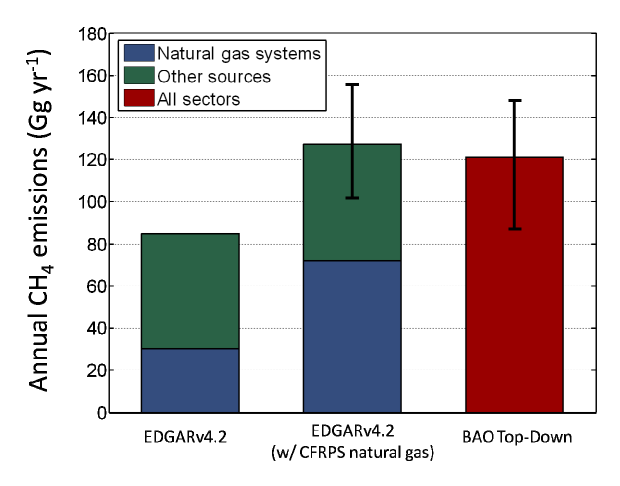


Figure 9 Annual CH₄ emissions for N/E wind sector for 2010. Figure shows a comparison of the EDGARv4.2 inventory, the EDGARv4.2 inventory using the CFRPS bottom-up estimate for the oil and gas sector, and the top-down estimate using observations at BAO. Error bar for the EDGARv4.2/CFRPS combined estimate reflects uncertainty in the CFRPS estimate. Error bar on the top-down estimate reflects the observed uncertainty, as propagated through Eq. (4), but does not include the uncertainty in the geographic area influencing observations at BAO. See Secs. 3.3.4 and 4.2 for a detailed discussion.